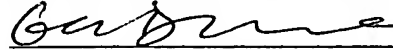


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Gaetano D. Maccarone

Registration No. 25,173

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**PATENT APPLICATION OF**

Jayprakash C. Bhatt

Brian D. Busch

Daniel P. Bybell

F. Richard Cottrell

Anemarie DeYoung

Chien Liu

Stephen J. Telfer

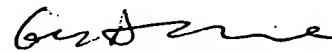
Jay E. Thornton

William T. Vetterling

FOR

THERMAL IMAGING SYSTEM

Respectfully submitted,



Gaetano D. Maccarone

Attorney for Applicants

Registration No. 25,173

POLAROID CORPORATION

Patent Department

784 Memorial Drive

Cambridge, MA 02139

Phone: 781-386-6405

Fax: 781-386-6435

THERMAL IMAGING SYSTEM

Reference To Related Applications

5 This application claims the benefit of prior provisional patent application serial no. 60/294,486, filed May 30, 2001 and prior provisional patent application serial no. 60/364,198, filed March 13, 2002.

Field Of The Invention

10 The present invention relates generally to a thermal imaging system and, more particularly, to a multicolor thermal imaging system wherein at least two image-forming layers of a thermal imaging member are addressed at least partially independently by a single thermal printhead or by multiple printheads from the same surface of the thermal imaging member.

Background of the Invention

15 Conventional methods for color thermal imaging such as thermal wax transfer printing and dye-diffusion thermal transfer typically involve the use of separate donor and receiver materials. The donor material typically has a colored image-forming material, or a color-forming imaging material, coated on a surface of a substrate and the image-forming material or the color-forming imaging material is
20 transferred thermally to the receiver material. In order to make multicolor images, a donor material with successive patches of differently-colored, or different color-forming, material may be used. In the case of printers having either interchangeable cassettes or more than one thermal head, different monochrome donor ribbons are utilized and multiple color separations are made and deposited successively above
25 one another. The use of donor members with multiple different color patches or the use of multiple donor members increases the complexity and the cost of such printing systems. It would be simpler to have a single-sheet imaging member that has the entire multicolor imaging reagent system embodied therein.

30 There have been described in the prior art numerous attempts to achieve multicolor, direct thermal printing. For example, there are known two-color direct thermal systems in which formation of the first color is affected by formation of the

second color. U.S. Patent 3,895,173 describes a dichromatic thermal recording paper which includes two leuco dye systems, one of which requires a higher activation temperature than the other. The higher temperature leuco dye system cannot be activated without activating the lower temperature leuco dye system.

5 There are known direct thermal imaging systems that utilize an imaging member having two color-forming layers coated on opposite surfaces of a transparent substrate. The imaging member is addressed by multiple printheads independently from each side of the imaging member. A thermal imaging system of this type is described in U.S. Patent 4,956,251.

10 Thermal systems that exploit a combination of dye transfer imaging and direct thermal imaging are also known. In systems of this type, a donor element and a receiver element are in contact with one another. The receiver element is capable of accepting dye, which is transferred from the donor element, and also includes a direct thermal color-forming layer. Following a first pass by a thermal printhead
15 during which dye is transferred from the donor element to the receiver element, the donor element is separated from the receiver and the receiver element is imaged a second time by a printhead to activate the direct thermal imaging material. This type of thermal system is described in U.S. Patent 4,328,977. U.S. Patent 5,284,816 describes a thermal imaging member that comprises a substrate having a direct
20 thermal color-forming layer on one side and a receiver element for dye transfer on the other side.

There are also known thermal imaging systems that utilize imaging members having spatially separated regions comprising direct thermal color-forming compositions that form different colors. U. S. Patents 5,618,063 and 5,644,352
25 describe thermal imaging systems in which different areas of a substrate are coated with formulations for forming two different colors. A similar bicolored material is described in U.S. Patent 4,627,641.

Another known thermal imaging system is a leuco-dye-containing, direct thermal system in which information is created by activating the imaging material at
30 one temperature and erased by heating the material to a different temperature. U.S. Patent 5,663,115 describes a system in which a transition from a crystalline to an

amorphous, or glass, phase is exploited to give a reversible color formation. Heating the imaging member to the melting point of a steroidal developer results in the formation of a colored amorphous phase while heating of this colored amorphous phase to a temperature lower than the crystalline melting point of the material causes
5 recrystallization of the developer and erasure of the image.

There is also known a thermal system containing one decolorizable, leuco dye containing, color-forming layer and a second leuco dye containing layer capable of forming a different color. The first color-forming layer colorizes at a low temperature while the second layer colorizes at a higher temperature, at which
10 temperature the decolorization of the first layer also takes place. In such systems, either one or the other color can be addressed at a particular point. U.S. Patent 4,020,232 discloses formation of one color by a leuco dye/base mechanism and the other by a leuco dye/acid mechanism wherein the color formed by one mechanism is neutralized by the reagent used to form the other. Variations of this type of system
15 are described in U.S. Patents 4,620,204; 5,710,094; 5,876,898 and 5,885,926.

Direct thermal imaging systems are known in which more than one layer may be addressed independently, and in which the most sensitive color-forming layer overlies the other color-forming layers. Following formation of an image in the layer outermost from the film base, the layer is deactivated by exposure to light
20 prior to forming images in the other, less sensitive, color-forming layers. Systems of this type are described in U.S. Patents 4,250,511; 4,734,704; 4,833,488; 4,840,933; 4,965,166; 5,055,373; 5,729,274; and 5,916,680.

As the state of the thermal imaging art advances and efforts are made to provide new thermal imaging systems that can meet new performance requirements, and to reduce or eliminate some of the undesirable requirements of the known
25 systems, it would be advantageous to have a muticolor thermal imaging system in which at least two different image-forming layers of a single imaging member can be addressed at least partially independently from the same surface by a single thermal printhead or by multiple thermal printheads so that each color can be printed
30 alone or in selectable proportion with the other color(s).

SUMMARY OF THE INVENTION

It is therefore an object of this invention to provide a multicolor thermal imaging system which allows for addressing, at least partially independently, with a single thermal printhead or multiple thermal printheads, at least two different image-forming layers of an imaging member from the same surface of the imaging member.

Another object of the invention is to provide such a multicolor thermal imaging system wherein each color can be printed alone or in selectable proportion with the other color(s).

Yet another object of the invention is to provide a multicolor thermal imaging system wherein at least two different image-forming layers of an imaging member are addressed at least partially independently by controlling the temperature applied to each of the layers and the time each of the layers is subjected to such temperature.

Still another object of the invention is to provide a multicolor thermal imaging system wherein at least two different image-forming layers of an imaging member are addressed at least partially independently with a thermal printhead or multiple thermal printheads from the same surface of the imaging member and one or more image-forming layers are addressed with a thermal printhead or multiple thermal printheads from the opposing surface of the imaging member.

A further object of the invention is to provide a multicolor thermal imaging system wherein at least two different image-forming layers of an imaging member are addressed at least partially independently with a single pass of a thermal printhead.

Another object of the invention is to provide a multicolor thermal imaging system which is capable of providing images which have adequate color separation for a particular application in which the system is used.

Still another object of the invention is to provide novel thermal imaging members.

These and other objects and advantages are accomplished in accordance with the invention by providing a multicolor thermal imaging system wherein at least

two, and preferably three, image-forming layers of a thermal imaging member can be addressed at least partially independently, from the same surface of the imaging member, by a single thermal printhead or by multiple thermal printheads. The advantageous thermal imaging system of the invention is based upon at least partially independently addressing a plurality of image-forming layers of a thermal imaging member utilizing two adjustable parameters, namely temperature and time. These parameters are adjusted in accordance with the invention to obtain the desired results in any particular instance by selecting the temperature of the thermal printhead and the period of time for which thermal energy is applied to each of the image-forming layers. According to the invention, each color of the multicolor imaging member can be printed alone or in selectable proportion with the other color(s). Thus, as will be described in detail, according to the invention the temperature-time domain is divided into regions corresponding to the different colors it is desired to combine in a final print.

The image-forming layers of the thermal imaging member undergo a change in color to provide the desired image in the imaging member. The change in color may be from colorless to a color or from colored to colorless or from one color to another color. The term "image-forming layer" as used throughout the application including in the claims, includes all such embodiments. In the case where the change in color is from colorless to a color, an image having different levels of optical density (i.e., different "gray levels") of that color may be obtained by varying the amount of color in each pixel of the image from a minimum density, D_{min} , which is substantially colorless, to a maximum density, D_{max} , in which the maximum amount of color is formed. In the case where the change in color is from colored to colorless, different gray levels are obtained by reducing the amount of color in a given pixel from D_{max} to D_{min} , where ideally D_{min} is substantially colorless. In this case, formation of the image involves converting a given pixel from a colored to a less colored, but not necessarily, colorless state.

A number of techniques can be used to achieve the advantageous results provided by exploiting the time and temperature variables in accordance with the invention. These include thermal diffusion with buried layers, chemical diffusion or

dissolution in conjunction with timing layers, melting transitions and chemical thresholds. Each of these techniques may be utilized alone, or in combination with others, to adjust the regions of the imaging member in which each desired color will be formed.

5 In a preferred embodiment, a thermal imaging member includes two, and preferably three, different image-forming layers carried by the same surface of a substrate. In another preferred embodiment, a thermal imaging member includes a layer or layers of image-forming material carried by one surface of a substrate and a layer or layers of image-forming material carried by the opposing surface of the
10 substrate. According to the imaging system of the invention, the image-forming layers of the imaging member can be addressed at least partially independently by a single thermal printhead or multiple printheads in contact with the same surface of the imaging member. In a preferred embodiment, one or two thermal printheads can be utilized to address at least partially independently from one surface of the
15 imaging member two different image-forming layers carried by one surface of the substrate and another thermal printhead utilized to address at least partially independently from the opposing surface of the imaging member one or more image-forming layers carried by the opposing surface of the substrate. The thermal printheads which contact the opposing surfaces of the imaging member can be
20 arranged directly opposite one another or offset from one another such that there is a delay between the times that any discrete area of the imaging member comes into contact with the respective thermal printheads.

 In another preferred embodiment one thermal printhead may be used to address at least partially independently two or more different image-forming layers
25 of the imaging member in a single pass and, optionally, a second thermal printhead used to address one or more image-forming layers, either in conjunction with the first thermal printhead, or subsequent thereto.

BRIEF DESCRIPTION OF THE DRAWINGS

 For a better understanding of the invention as well as other objects and
30 advantages and further features thereof, reference is made to the following detailed

description of various preferred embodiments thereof taken in conjunction with the accompanying drawings wherein:

Fig. 1 is a graphical representation of the colors which may be printed by a prior art two-color, direct thermal printing system;

5 Fig. 2 is a graphical representation of the colors which may be printed by a two-color direct thermal printing embodiment of the invention;

Fig. 3 is a graphical illustration of non-independent colored-dot formation encountered in prior art direct thermal printing;

10 Fig. 4 is a graphical representation of the colors which may be printed by a prior art three-color direct thermal printing system and by a three-color direct thermal printing embodiment of the invention;

Fig. 5 is a graphical representation illustrating one embodiment of the invention;

15 Fig. 6 is a graphical representation further illustrating the embodiment of the invention illustrated in Fig. 5;

Fig. 7 is a graphical representation illustrating the practice of a three-color embodiment of the invention;

Fig. 8 is a partially schematic, side sectional view of a two color imaging member according to the invention which utilizes thermal delays;

20 Fig. 9 is a partially schematic, side sectional view of a three color imaging member according to the invention which utilizes thermal delays;

Fig. 10 is a partially schematic, side sectional view of another three color imaging member according to the invention which utilizes thermal delays;

25 Fig. 11 is a partially schematic, side sectional view of a thermal printing apparatus for carrying out an embodiment of the invention:

Fig. 12 is a graphical representation of a method for applying voltage to a conventional thermal printhead during a prior art thermal imaging method;

30 Fig. 13 is a graphical representation of a method for applying voltage to a conventional thermal printhead in the practice of an embodiment of the thermal imaging system of the invention;

Fig. 14 is a graphical representation of another method for applying voltage to a conventional thermal printhead in the practice of an embodiment of the thermal imaging system of the invention;

5 Fig. 15 is a graphical representation showing the development time of two dyes as a function of temperature;

Fig. 16 is a partially schematic, side sectional view of a multicolor imaging member according to the invention which utilizes chemical diffusion and dissolution;

10 Fig. 17 is a partially schematic, side sectional view of a negative-working multicolor imaging member according to the invention; and

Fig. 18 is a partially schematic, side sectional view of a three color imaging member according to the invention which utilizes chemical diffusion and dissolution.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

15 As previously mentioned, according to the multicolor thermal imaging system of the invention, two or more image-forming layers of a multicolor thermal imaging member are addressed at least partially independently from the same surface of the imaging member, so that each color may be printed alone or in selectable proportion with the others, and these results are accomplished by selecting
20 the colors on the basis of two adjustable parameters, namely temperature and time. The temperature – time domain is divided into regions corresponding to the different colors it is desired to combine.

To assist those skilled in the art to better understand the concept of independent control of color, as applied to multicolor direct thermal printing
25 according to the present invention, it is helpful to consider first a prior art thermal imaging system involving a thermal imaging member containing two color-forming layers on a white reflective substrate. For the purpose of discussion it will be considered that one layer is a cyan color-forming layer and the other a magenta color-forming layer and, further, that the cyan layer has a temperature threshold
30 above that of the magenta layer. If a fixed-length thermal pulse is applied to a

discrete point, or area, on this imaging member, a color will form depending upon the magnitude of the pulse. Pulses of increasing magnitude lead to increasing peak temperature in the image-forming layers at the location of the thermal pulse. The originally white medium will become progressively more magenta as the magenta threshold temperature for coloration is exceeded and then progressively more blue, i.e., magenta plus cyan, as the cyan threshold temperature for coloration is exceeded. This progression of color may be represented by the two-dimensional color diagram illustrated in Fig. 1.

As shown by the curvilinear path, the color first moves in the magenta direction as the threshold temperature is exceeded in the magenta layer and then in the cyan direction, i.e., towards blue, as the threshold temperature is surpassed in the cyan layer. Each point on the color path is associated with the magnitude of the thermal pulse that created it and there is a fixed ratio of magenta and cyan color associated with each pulse magnitude. A similar progression of colors is produced if the applied pulse has a fixed magnitude and variable duration provided that the power is sufficient ultimately to raise both dye layers above their threshold coloration temperatures. In this case, when the pulse begins, the two dye layers will advance in temperature. For longer and longer pulse durations the dye temperatures will first exceed the magenta threshold and then the cyan threshold. Each pulse duration will correspond to a well-defined color, again passing from white to magenta to blue along a curvilinear path. Prior art thermal imaging systems, using either a modulation of pulse amplitude or pulse duration, are therefore essentially limited to the reproduction of colors falling on curvilinear paths in the color space.

The present invention, by addressing at least partially independently the different image-forming layers of a multicolor thermal imaging member, provides a thermal imaging method in which the colors formed are not constrained by a one dimensional path but can instead be selected throughout regions on both sides of the path as is illustrated in the shaded region of Fig. 2.

In the foregoing description the term "partially independently" is used to describe the addressing of the image-forming layers. The degree to which the image-forming layers can be addressed independently is related to the image

property commonly referred to as "color separation". As stated previously, an object of the invention is to provide images with adequate color separation for the various applications for which the present thermal imaging method is suitable. For example, photographic imaging requires that the color separation be comparable to that which can be obtained with conventional photographic exposure and development. Depending upon the printing time, available printing power, and other factors, various degrees of independence in the addressing of the image-forming layers can be achieved. The term "independently" shall be used to refer to instances in which the printing of one color-forming layer typically results in a very small, but not generally visible optical density (density < 0.05) in the other color-forming layer(s). In the same manner, the term "substantially independent" color printing will be used to refer to instances in which inadvertent or unintentional coloration of another image-forming layer or layers results in a visible density which is at a level typical of interimage coloration in multicolor photography (density < 0.2). In some instances color crosstalk at this level is considered photographically desirable. The term "partially independent" addressing of the image-forming layers is used to refer to instances in which the printing of maximum density in the layer being addressed results in the coloration of another image-forming layer or layers at a density higher than 0.2 but not higher than about 1.0. The phrase "at least partially independently" is inclusive of all of the degrees of independence described above.

A distinction between the thermal imaging system of the invention and the prior art thermal imaging methods can be seen from the nature of the images which are obtainable from each. When two image-forming layers are not addressable independently one or both of them will not be able to be printed without substantial color contamination from the other. For example, consider a single-sheet thermal imaging member which is designed to provide two colors, Color 1 and Color 2, with temperature thresholds for coloration of, respectively, T_1 and T_2 where $T_1 > T_2$. Consider the attempt to form a dot of a single color using a heating element to heat the thermal member from the top surface. There will be a point, typically in the center of the heated area, where the temperature T takes its highest value, T_{max} . Away from this point T is lower, falling off quickly outside of the heated area to a

temperature well below T_1 or T_2 , as indicated schematically in Fig. 3a. A "clean" dot of Color 2 may be printed in regions where the local temperature T is greater than T_2 but less than T_1 (see Fig. 3b). If T_{max} exceeds T_1 , then the dot will be contaminated with Color 1 in the center and independent color formation will no longer be possible.

It is notable that an attempt to print a dot of Color 1 will require that $T_{max} > T_1$, and since $T_1 > T_2$ this will inevitably mean that Color 2 will be printed as well (see Fig. 3c). Consequently, independent printing of Color 1 is not possible. An attempt can be made to correct this problem by incorporating a bleaching of Color 2 which occurs whenever Color 1 is formed. If bleaching is carried out, only Color 1 will be visible in the heated region where T is greater than T_1 . However, this does not constitute independent addressing for two reasons. First, it is not possible to obtain arbitrary mixtures of Color 1 and Color 2 in this manner. Second, there remains an annular region around each dot of Color 1 within which Color 2 is not bleached (see Fig. 3d).

According to the present invention, independent addressing of both colors in the above example is achieved by introducing a timing mechanism by which the coloration of the second dye layer is delayed with respect to the coloration of the first dye layer. During this delay period, it is possible to write on the first dye layer without colorizing the second; and, if the second layer has a lower threshold temperature for coloration than the first, it will later be possible to write on the second without exceeding the threshold of the first.

In one embodiment, the method of the invention will allow completely independent formation of cyan or magenta. Thus, in this embodiment, one combination of temperature and time will permit the selection of any density of magenta on the white-magenta axis while not producing any noticeable cyan color. Another combination of temperature and time will permit the selection of any density of cyan on the white-cyan axis while not producing any noticeable magenta coloration. A juxtaposition of two temperature-time combinations will allow the selection of any cyan/magenta mixture within the enclosed area indicated on Fig. 2, thus providing independent control of cyan and magenta.

In other embodiments of the invention, thermal addressing of the image-forming layers, rather than being completely independent, can be substantially independent or only partially independent. Various considerations, including material properties, printing speed, energy consumption, material costs and other system requirements may dictate a system with increased color cross-talk. While independent or substantially independent color selection according to the invention is desirable for photographic-quality printing, this requirement is of less importance in the printing of certain images such as, for example, product labels or multicolor coupons, and in these instances may be sacrificed for economic considerations such as improved printing speed or lower costs.

In these embodiments of the invention where addressing of the separate image-forming layers of a multicolor thermal imaging member is not completely, but rather substantially, or partially, independent, and by design the printing of cyan may produce a controlled amount of magenta color formation and vice-versa, it will not be possible to print completely pure magenta or completely pure cyan. Indeed, there will be a region of the color box near each coordinate axis that represents unprintable colors and the available colors will fall into a more restricted region such as the shaded area illustrated schematically in Fig. 2. In these instances, although the palette of colors available is less than the selection encompassed by the embodiments of the invention where color selection is controlled completely independently, it is nevertheless greatly superior to the very restricted selection of colors allowed by the prior art systems.

Similar considerations apply to three-color embodiments of the present invention. For these embodiments, the color space is three-dimensional and is commonly referred to as a "color cube" as is illustrated in Fig. 4. If fixed-length thermal pulses of increasing temperature are applied to a prior art multicolor direct thermal printing medium, it is possible to produce colors which fall on a curvilinear path through the cube as illustrated by the dotted arrow. As seen, the path extends from one color, usually white, to another color, usually black, while passing through a fixed variety of colors. In comparison, one embodiment of the present invention advantageously provides the capability to print any color within the three-

dimensional color cube. In other embodiments of the invention, where addressing of the color-forming layers is substantially or partially independent, formation of colors within the shaded area of Fig. 4 is possible, again providing considerably more flexibility in the choice of colors than that offered by prior art direct thermal printing systems.

For the purpose of describing the temperature and time parameter feature of the invention, reference is made to Fig. 5 which is a graphical representation of one embodiment of the invention. For example, the thermal imaging member may contain a cyan image-forming material which provides a visible cyan color region, C, when subjected to a relatively high temperature for a short period of time and a magenta image-forming material which provides a visible magenta region, A, when subjected to a lower temperature for a longer period of time. A combination of short and long pulses of heat at different temperatures can be utilized to select the proportions of each color. It can be seen that according to the invention, since there are two adjustable variables involved and two or more image-forming materials, at least substantially complete independent control of any particular color according to the invention requires that each color be assigned a substantially unique range of time and temperature.

Other considerations relevant to the multicolor thermal imaging system of the invention can be understood from the following discussion of a two-color leuco dye system in conjunction with Fig. 6. Consider, for example, a system wherein color is generated by a leuco dye that is being thermally diffused to combine with an acid developer material. In this instance, it may not be possible to constrain the colorant response to a completely enclosed region such as is shown in Fig. 5.

Although it may be intended to utilize temperatures and time periods within the regions shown in Fig. 5 the imaging member may also be responsive at a wider range of temperatures and time periods. Referring now to Fig. 6 it can be seen that in this illustrative example, regions A and C would be the regions selected for printing magenta and cyan, respectively. However, the temperature and time combinations in regions B and E, for example, will also be adequate to permit diffusion of the magenta leuco dye to the developer. Also, cyan will be printed for temperature –

time combinations in regions D and E. Thus, in order to obtain substantially complete independent control of cyan and magenta image-forming materials according to the invention a magenta printing region, A, should preferably be selected such that it does not overlap regions C, D or E, or any other region in which cyan is responsive. Conversely, cyan printing region, C, should preferably be selected such that it does not overlap regions A, B and E, or any other region in which magenta is responsive. Generally, this means that for the illustrative diffusive leuco dye system, the separately selected color printing regions should be arranged along a slope decreasing from higher to lower time periods and from lower to higher temperatures. It will be appreciated that in actual implementations, the chosen printing regions may not be rectangular in shape as shown in the schematic representation, but will have a shape governed by the behavior of the physical process that leads to coloration, and may contain limited regional overlap consistent with the desired color separation for a particular application.

A suitable schematic arrangement for a three-color diffusion-controlled leuco dye system according to the invention is illustrated in Fig. 7 where the time - temperature combinations for printing magenta, cyan and yellow, respectively, are shown.

In preferred embodiments of the invention, the temperatures selected for the color-forming regions generally are in the range of from about 50°C to about 450°C. The time period for which the thermal energy is applied to the color-forming layers of the imaging member is preferably in the range of from about 0.01 to about 100 milliseconds.

As mentioned previously, a number of image-forming techniques may be exploited in accordance with the invention including thermal diffusion with buried layers, chemical diffusion or dissolution in conjunction with timing layers, melting transitions and chemical thresholds.

Referring now to Fig. 8 there is seen a multicolor thermal imaging member that utilizes thermal time delays to define the printing regions for the respective colors to be formed. The imaging member 10 relies upon the diffusion of heat through the imaging member in order to obtain the timing differences that are

exploited according to the invention. Imaging member 10 includes a substrate 12 carrying cyan and magenta image-forming layers, 14 and 16, respectively, and spacer interlayer 18. It should be noted here that in various embodiments of the invention the image-forming layers may themselves comprise two or more separate
5 layers. For example, where the image-forming material is a leuco dye which is used in conjunction with a developer material, the leuco dye and developer material may be disposed in separate layers.

Where the imaging member 10 is heated by a thermal printhead from above cyan image-forming layer 14 the heat will penetrate into the imaging member to
10 reach magenta image-forming layer 16. Cyan image-forming layer 14 will be heated above its coloration threshold temperature almost immediately by the thermal printhead after the heat is applied, but there will be a more significant delay before the magenta image-forming layer 16 approaches its threshold temperature. If both image-forming layers were such as to begin forming color at the same temperature,
15 e.g., 120°C, and the printhead were to heat the surface of imaging member 10 to a temperature of substantially more than 120°C, then the cyan image-forming layer 14 would begin to provide cyan color almost at once whereas magenta image-forming layer 16 would begin to provide magenta color after a time delay dependent upon the thickness of spacer layer 18. The chemical nature of the activation of the color
20 in each layer would not be critical.

To provide multicolor printing in accordance with the invention each image-forming layer is arranged to be activated at a different temperature, e.g., T_5 for cyan image-forming layer 14 and T_6 for the "buried" magenta image-forming layer 16. This result can be achieved, for example, by arranging these image-forming layers to
25 have different melting temperatures or by incorporating in them different thermal solvents, which will melt at different temperatures and liquefy the image-forming materials. Temperature T_5 is selected to be higher than T_6 .

Where a temperature less than T_6 is applied to the imaging member for any length of time no color will be formed. Thus, the imaging material may be shipped
30 and stored safely at a temperature less than T_6 . Where a printing element in contact with layer 14 applies such heating as to cause a temperature between T_5 and T_6 to be

attained by image-forming layer 16, then the cyan image-forming layer 14 will remain substantially colorless and magenta image-forming layer 16 will develop magenta color density after a time delay which is a function of the thickness of spacer layer 18. Where a temperature just above T_5 is applied to the imaging member by a printing element in contact with image-forming layer 14, then the cyan image-forming layer 14 will begin developing color density immediately and magenta image-forming layer 16 will also develop magenta color density but only after a time delay. Said another way, at intermediate temperatures and relatively long time periods it is possible to produce magenta color without cyan color and for high temperatures and relatively short time periods, it is possible to produce cyan color without any magenta color. A relatively short, high temperature heat pulse juxtaposed with a longer, intermediate temperature heat pulse will result in the combination of magenta and cyan colors in selected proportions.

It will be appreciated by those skilled in the art that the mechanisms described above in reference to Fig. 8 will provide optimum differentiation between the two colors where the thermal printhead is chosen so as to conduct heat away efficiently from the surface of imaging member 10 after the application of heat. This is particularly important immediately following printing a pixel in image-forming layer 14.

The image-forming layers 14 and 16 of imaging member 10 may optionally undergo more than one color change. For example, image-forming layer 14 may go from colorless to yellow to red as a function of the heat applied. Image-forming layer 16 could initially be colored, then become colorless and then go to a different color. Those skilled in the art will recognize that such color changes can be obtained by exploiting the imaging mechanism described in U.S. Patent 3,895,173.

Any known printing modality may be used to provide a third image-forming layer or additional image-forming layers beyond the two illustrated in Fig. 8. For example, the third image-forming layer may be imaged by ink jet printing, thermal transfer, electrophotography, etc. In particular, imaging member 10 may include a third image-forming layer which, after color is formed in the layer, can then be fixed by exposure to light as is known in the art. In this embodiment, the third image-

forming layer should be positioned close to the surface of imaging member 10 and printed at a lower temperature than image-forming layer 14, prior to the printing of image-forming layer 14. Fixation of this third layer should also occur prior to printing of image-forming layer 14.

5 Substrate 12 may be of any suitable material for use in thermal imaging members, such as polymeric materials, and may be transparent or reflective.

Any combination of materials that may be thermally induced to change color may be used. The materials may react chemically under the influence of heat, either as a result of being brought together by a physical mechanism, such as melting or
10 diffusion, or through thermal acceleration of a reaction rate. The reaction may be chemically reversible or irreversible.

For example, a colorless dye precursor may form color upon heat-induced contact with a reagent. This reagent may be a Bronsted acid, as described in "Imaging Processes and Materials", Neblette's Eighth Edition, J. Sturge, V.
15 Walworth, A. Shepp, Eds., Van Nostrand Reinhold, 1989, pp. 274-275, or a Lewis acid, as described for example in U.S. Patent No. 4,636,819. Suitable dye precursors for use with acidic reagents are described, for example, in U.S. Patent No. 2,417,897, South African Patent 68-00170, South African Patent 68-00323 and Ger. Offen. 2,259,409. Further examples of such dyes may be found in "Synthesis and
20 Properties of Phthalide-type Color Formers", by Ina Fletcher and Rudolf Zink, in "Chemistry and Applications of Leuco Dyes", Muthyala Ed., Plenum Press, New York, 1997. Such dyes may comprise a triarylmethane, diphenylmethane, xanthene, thiazine or spiro compound, for example, Crystal Violet Lactone, N-halophenyl leuco Auramine, rhodamine B anilinolactam, 3-piperidino-6-methyl-7-anilino-
25 fluoran, benzoyl leuco Methylene blue, 3-methyl-spirodinaphthofuran, etc. The acidic material may be a phenol derivative or an aromatic carboxylic acid derivative, for example, p-tert-butylphenol, 2,2-bis (p-hydroxyphenyl)propane, 1,1-bis(p-hydroxyphenyl) pentane, p-hydroxybenzoic acid, 3,5-di-tert-butylsalicylic acid, etc. Such thermal imaging materials and various combinations thereof are now
30 well known, and various methods of preparing heat-sensitive recording elements

employing these materials also are well known and have been described, for example, in U.S. Patents Nos. 3,539,375, 4,401,717 and 4,415,633.

The reagent used to form a colored dye from a colorless precursor may also be an electrophile, as described, for example, in U.S. Patent No. 4,745,046, a base, as described, for example, in U.S. Patent No. 4,020,232, an oxidizing agent, as
5 described, for example, in U.S. Patents Nos. 3,390,994 and 3,647,467, a reducing agent, as described, for example, in U.S. Patent No. 4,042,392, a chelatable agent, as described, for example, in U.S. Patent No. 3,293,055 for spiropyran dyes, or a metal ion, as described, for example, in U.S. Patent No. 5,196,297 in which thiolactone
10 dyes form a complex with a silver salt to produce a colored species.

The reverse reaction, in which a colored material is rendered colorless by the action of a reagent, may also be used. Thus, for example, a protonated indicator dye may be rendered colorless by the action of a base, or a preformed dye may be irreversibly decolorized by the action of a base, as described, for example, in U.S.
15 Patents Nos. 4,290,951 and 4,290,955, or an electrophilic dye may be bleached by the action of a nucleophile, as described in U.S. Patent No. 5,258,274.

Reactions such as those described above may also be used to convert a molecule from one colored form to another form having a different color.

The reagents used in schemes such as those described above may be sequestered from the dye precursor and brought into contact with the dye precursor
20 by the action of heat, or alternatively a chemical precursor to the reagents themselves may be used. The precursor to the reagent may be in intimate contact with the dye precursor. The action of heat may be used to release the reagent from the reagent precursor. Thus, for example, U.S. Patent No. 5,401,619 describes the
25 thermal release of a Bronsted acid from a precursor molecule. Other examples of thermally-releasable reagents may be found in "Chemical Triggering", G. J. Sabongi, Plenum Press, New York (1987).

Two materials that couple together to form a new colored molecule may be employed. Such materials include diazonium salts with appropriate couplers, as
30 described, for example, in "Imaging Processes and Materials" pp. 268-270 and U.S. Patent No. 6,197,725, or oxidized phenylenediamine compounds with appropriate

couplers, as described, for example, in U.S. Patents Nos. 2,967,784, 2,995,465, 2,995,466, 3,076,721, and 3,129,101.

Yet another chemical color change method involves a unimolecular reaction, which may form color from a colorless precursor, cause a change in the color of a colored material, or bleach a colored material. The rate of such a reaction may be accelerated by heat. For example, U.S. Pat. No. 3,488,705 discloses thermally unstable organic acid salts of triarylmethane dyes that are decomposed and bleached upon heating. U.S. Pat. No. 3,745,009 reissued as U.S. Pat. No. Re. 29,168 and U.S. Pat. No. 3,832,212 disclose heat-sensitive compounds for thermography containing a heterocyclic nitrogen atom substituted with an -OR group, for example, a carbonate group, that decolorizes by undergoing homolytic or heterolytic cleavage of the nitrogen-oxygen bond upon heating to produce an RO⁺ ion or RO' radical and a dye base or dye radical which may in part fragment further. U.S. Pat. No. 4,380,629 discloses styryl-like compounds which undergo coloration or bleaching, reversibly or irreversibly via ring-opening and ring-closing in response to activating energies. U.S. Patent No. 4,720,449 describes an intramolecular acylation reaction which converts a colorless molecule to a colored form. U.S. Patent No. 4,243,052 describes a pyrolysis of a mixed carbonate of a quinophthalone precursor which may be used to form a dye. U.S. Patent No. 4,602,263 describes a thermally-removable protecting group which may be used to reveal a dye or to change the color of a dye. U.S. Patent No. 5,350,870 describes an intramolecular acylation reaction which may be used to induce a color change. A further example of a unimolecular color-forming reaction is described in "New Thermo-Response Dyes: Coloration by the Claisen Rearrangement and Intramolecular Acid-Base Reaction Masahiko Inouye, Kikuo Tsuchiya, and Teijiro Kitao, *Angew. Chem. Int. Ed. Engl.* **31**, pp. 204-5 (1992).

It is not necessary that the colored material formed be a dye. The colored species may also be, for example, a species such as a metal or a polymer. U.S. Patent No. 3,107,174 describes the thermal formation of metallic silver (which appears black) through reduction of a colorless silver behenate salt by a suitable reducing

agent. U.S. Patent No. 4,242,440 describes a thermally-activated system in which a polyacetylene is used as the chromophore.

Physical mechanisms may also be used. Phase changes leading to changes in physical appearance are well known. The phase change may for example lead to a change in scattering of light. Thermally-activated diffusion of dye from a restricted area, thereby changing its covering power and apparent density, has also been described in "A New Thermographic Process", by Shoichiro Hoshino, Akira Kato, and Yuzo Ando, Symposium on Unconventional Photographic System, Washington D.C. October 29, 1964.

Image-forming layers 14 and 16 may comprise any of the image-forming materials described above, or any other thermally-activated colorants, and are typically from about 0.5 to about 4.0 μm in thickness, preferably about 2 μm . In the case where image-forming layers 14 and 16 comprise more than one layer, each of the constituent layers are typically from about 0.1 to about 3.0 μm in thickness.

Image-forming layers 14 and 16 may comprise dispersions of solid materials, encapsulated liquid, amorphous or solid materials or solutions of active materials in polymeric binders, or any combinations of the above.

Interlayer 18 is typically from about 5 to about 30 μm in thickness, preferably about 14 - 25 μm . Interlayer 18 may comprise any suitable material including inert materials or materials which undergo a phase change upon heating such as where the layer includes a thermal solvent. Typical suitable materials include polymeric materials such as poly (vinyl alcohol). Interlayer 18 may comprise one or more suitable materials and can be made up of one or more layers.

Interlayer 18 can be coated from aqueous or solvent solution or applied as a film laminated to the image-forming layers. Interlayer 18 can be opaque or transparent. Where the interlayer is opaque, substrate 12 is preferably transparent so either outer surface of imaging member 10 can be printed with a thermal printhead from one side. In a particularly preferred embodiment, substrate 12 is transparent and interlayer 18 is white. The effect of two-sided printing of a single sheet using only a single thermal printhead, printing on only one side of said sheet, is thereby obtained.

The thermal imaging members of the invention may also include thermal backcoat layers and protective topcoat layers arranged over the outer surface of the image-forming layers. In a preferred embodiment of the imaging member shown in Fig. 8, there are included a barrier coating and a protective topcoat layer over layer 14. The barrier layer may comprise water and gas inhibiting materials. Taken together, the barrier and topcoat layers may provide protection from UV radiation.

In an alternative embodiment of the imaging member shown in Fig. 8, image-forming layer 16 is coated on a thin substrate 12 such as, for example, poly(ethylene terephthalate) having a thickness of about 4.5 μm . Interlayer 18 and image-forming layer 14 are then deposited. Substrate 12 may be opaque or transparent and can be coated, laminated or extruded onto layer 16. In this embodiment of the invention, image-forming layers 14 and 16 can be addressed by a thermal printhead or printheads through the thin substrate 12.

Referring now to Fig. 9 there is seen a three color thermal imaging member according to the invention that utilizes thermal delays to define the printing regions for the colors to be formed. The three color imaging member 20 includes substrate 22, cyan, magenta and yellow image-forming layers, 24, 26 and 28, respectively, and spacer interlayers 30 and 32. Preferably, interlayer 30 is thinner than interlayer 32 so long as the materials comprising both layers have the same heat capacity and thermal conductivity. The activation temperature of layer 24 is higher than that of layer 26 which in turn is higher than that of layer 28.

According to a preferred embodiment of the invention a thermal imaging member in which a plurality of image-forming layers are carried by the same surface of a substrate, as is illustrated in Fig. 9 where three image-forming layers are carried by the same surface of substrate 22, two of the image-forming layers can be imaged by one or more thermal printheads from one surface of the member and at least a third image-forming layer imaged by a separate thermal printhead from the opposite side of the substrate. In the embodiment illustrated in Fig. 9, image-forming layers 24 and 26 are imaged by one or two thermal printheads in contact with the outer surface of color-forming layer 24 and color-forming layer 28 is imaged by a thermal printhead in contact with the outer surface of substrate 22. In this embodiment of

the invention, substrate 22 is relatively thin and is typically less than about 20 μm and preferably about 5 μm thick.

In this instance, since the substrate 22 is relatively thin, it is preferred to laminate the imaged member to another base such as label card stock material. Such
5 laminate structures can also provide additional features such as where the image-forming layers are designed to separate when the laminated structure is taken apart, thus providing security features. Also, ultraviolet and infrared security features can be incorporated into the image-forming layers.

By laminating the imaged thermal imaging member to another base, a
10 number of product applications are provided. The base stock can be anything that will support an adhesive bonding agent. Thus, imaging can be carried out on various materials such as transparent or reflective sticker materials which can be laminated onto transparent or reflective carrier materials to provide transparencies or reflective products.

15 Fig. 10 illustrates a multicolor thermal imaging member according to the invention wherein two image-forming layers are arranged on one side of a substrate and one image-forming layer is arranged on the other side of the substrate. Referring now to Fig. 10 there is seen imaging member 40 which includes a substrate 42, a first image-forming layer 44, interlayer 46, a second image-forming
20 layer 48, a third image-forming layer 50, an optional white or reflective layer 52, a backcoat layer 53 and a topcoat layer 54. In this preferred embodiment substrate 42 is transparent. The image-forming layers and the interlayer may comprise any of the materials described above for such layers. Optional layer 52 may be any suitable reflective material or may comprise particles of a white pigment such as titanium
25 dioxide. Protective topcoat and backcoat layers 53 and 54 may comprise any suitable materials providing the functions of lubrication, heat resistance, UV, water and oxygen barrier properties, etc. Such materials may comprise polymeric binders in which appropriate small molecules are dissolved or dispersed, as will be familiar to those skilled in the art. The activation temperature of image-forming layer 48 is
30 lower than that of image-forming layer 44 and the activation temperature of image-forming layer 50 can be the same as that of image-forming layer 48 or higher or

lower and may be as low as possible consistent with the requirement of room temperature and shipping stability.

In a preferred embodiment, one thermal printhead can be utilized to address independently from one surface of the imaging member two image-forming layers
5 carried by one surface of a substrate and another thermal printhead utilized to address independently from the opposing surface of the imaging member one or more image-forming layers carried by the opposing surface of the substrate. This preferred embodiment of the invention will be described further in detail with respect to the imaging member shown in Fig. 10 although it will be understood that
10 the embodiment may be practiced with other suitable imaging members. The thermal printheads which are brought into contact with opposing surfaces of the imaging member can be arranged directly opposite to each other. Alternatively, and preferably, the respective printheads are offset from each other as is illustrated in Fig. 11. Further, two separate thermal print engines such as an Alps MBL 25,
15 available from Alps Electric Co. Ltd., Tokyo, Japan can be used. However, it is preferred to utilize a thermal printing apparatus where some of the components such as the drive motor and power source are shared by the two print stations.

Referring now to Fig. 11 there is seen a roll of a thermal imaging member 55, for example, the imaging member illustrated in Fig. 10. The imaging member is
20 passed between a first thermal printhead 56 and backing roller 57 and subsequently between a second thermal printhead 58 and backing roller 59. First thermal printhead 56 addresses at least partially independently the first and second image-forming layers 44 and 48, which may be cyan and magenta image-forming layers respectively and second thermal printhead 58 addresses third image-forming layer
25 50 which may be a yellow image-forming layer.

As discussed previously, in the advantageous multicolor thermal imaging method of the invention, two or more different image-forming layers of a thermal imaging member are addressed at least partially independently from the same surface of the imaging member by a single thermal printhead or multiple thermal
30 printheads. In a particularly preferred embodiment of the invention, two or more different image-forming layers of a thermal imaging member are addressed at least

partially independently by a single thermal printhead in a single pass. The methods for doing so can be carried out by the manipulation of control signals applied to a conventional thermal printhead, the heating elements of which are in contact with a surface of the imaging member. A conventional thermal printhead is composed of a linear array of heating elements, each having a corresponding electronic switch
5 capable of connecting it between a common voltage bus and ground. The voltage of the common bus and the time that the electrical switch is closed will together affect the temperature and time of the thermal exposure.

In order to describe the methods for controlling temperature in the practice of the invention, the operation of the thermal printhead will now be described in more
10 detail. In normal use of the printhead, a fixed voltage is applied to the printhead and the modulation of density on the image formed is achieved by controlling the length of time that power is applied to the heating elements. The control system may be discrete, that is, the time interval used to print each pixel on the imaging member is divided into a number of discrete subintervals and the heating element may be either
15 active or inactive during each of the subintervals. Moreover, the duty cycle of the heating within each subinterval may be controlled. For example, if a heating element is active during one of the subintervals and the duty cycle for that subinterval is 50%, then power will be applied to the heating element during 50% of that particular subinterval. This process is illustrated in Fig. 12.

Fig. 12 illustrates a printhead application in which each pixel-printing interval is divided into seven equal subintervals. For the case illustrated, the pixel is active for the first four subintervals and then inactive for three subintervals. In addition, the voltage pulses that are applied have a 50% duty cycle, so that within
25 each active subinterval, the voltage is on for half of the subinterval and off for the other half. Insofar as the temperature of the heating element is responsive to the power applied, it is easily appreciated by those skilled in the art that this temperature may be affected by the common bus voltage and by the duty cycle of the pulses. In fact, if the individual subintervals are much shorter than the thermal time constant
30 for heating and cooling of the medium, then the effect of changing the voltage of the

common bus may be mimicked by the effect of changing the duty cycle of the pulses.

5 This offers at least two possibilities for controlling the average power applied to the printhead. The first is that the temperature of a printhead heating element may be controlled by manipulating the voltage on the common bus, while the duty cycle remains fixed at some predetermined values for each subinterval. In this instance, the temperature is controlled primarily by the choice of bus voltage, and the time is controlled by the selection of the number of subintervals for which the heater is activated.

10 The second possibility is the control of the heater temperature by manipulation of the duty cycles of the subintervals while the bus voltage remains fixed. Best use of this method of temperature control requires that the subintervals be short compared to the thermal time-constant of the imaging member, so that the temperature in the image-forming layer responds to the average power applied during the subinterval rather than tracking the rapid voltage transitions. For a typical printhead in this application, the subinterval time may be ten or more times shorter than the thermal response time of the imaging member so this condition is well satisfied.

20 The choice between these two methods of control, or of a combination of the two, is a matter of practical design. For example, in a multiple-pass system in which each color layer is printed in a separate pass of the imaging member beneath the printhead, it is not difficult to change the voltage applied to the printhead common bus on each pass. The applied voltages can then be easily adjusted for best results. On the other hand, for a single-pass system in which two or more color layers are written in quick succession at each pixel, it is generally more convenient and economical to operate the head at a fixed voltage. In this case the temperature changes are preferably effected by a predetermined sequence of duty cycles of the subintervals.

25 The two techniques are illustrated in Figs. 13 and 14 which are based on a two image-forming layer system in which one image-forming layer is activated by a

30

high temperature applied for short times, and the other image-forming layer is activated by a lower temperature applied for longer times.

Fig. 13 illustrates schematically a method of alternately writing on the two image-forming layers by changing the bus voltage and the time over which the heater is activated. Initially the writing is at high-temperature for a short time, and is accomplished by a short series of high voltage pulses. Subsequently, writing is done at a low temperature for a long time by using a longer sequence of lower-voltage pulses. The sequence then repeats to alternate back and forth between color-forming layers.

Fig. 14 illustrates schematically another method of alternately writing on two image-forming layers. In this case the pulse duty cycle is varied rather than the pulse voltages. The high-temperature, short-time heating is performed with a short sequence of pulses having a large duty cycle. The low-temperature, long-time heating is performed with a longer sequence of pulses having a low duty cycle.

The method illustrated in Figure 14 for forming an image in an imaging member of the invention with two image-forming layers will now be described in more detail. The time interval for forming a single pixel of an image in the region of the thermal imaging member that is in thermal contact with a heating element of the printhead is divided into a plurality of temporal subintervals (hereinafter referred to as mini-subintervals), as described above. The mini-subintervals may be equal or different in duration to each other. In a preferred embodiment, the mini-subintervals are of equal duration. The time interval for forming a single pixel is also divided into a first and a second time interval, the first time interval being shorter than the second time interval. The first time interval is used to form an image in a first color-forming layer of the thermal imaging member (which may be a higher-temperature color-forming layer), and the second time interval is used to form an image in a second color-forming layer of the thermal imaging member (which may be a lower-temperature color-forming layer). The first time interval and the second time interval will, between them, contain most or all of the mini-subintervals described above. In the case when the mini-subintervals are of equal duration, the first time interval will contain fewer mini-subintervals than the second time interval. It is

preferred that the second time interval be at least twice as long as the first time interval. It is not necessary that the first time interval precede the second time interval. It is possible that, in combination, the first time interval and the second time interval do not occupy the entire time interval for printing a single pixel.

5 However, it is preferred that, in combination, the first time interval and the second time interval occupy most of the time interval for printing a single pixel.

A heating element of the printhead is activated by applying a single pulse of electrical current during a mini-subinterval. The proportion of the duration of the mini-subinterval (i.e., the duty cycle) during which this pulse of electrical current is
10 applied may take any value between about 1% and 100%. In a preferred embodiment, the duty cycle is a fixed value, p_1 , during the first time interval, and a second fixed value, p_2 , during the second time interval, and $p_1 > p_2$. In a preferred embodiment, p_1 approaches 100%. It is preferred that p_1 be greater than or equal to twice the length of p_2 .

15 Within the first time interval and the second time interval, different degrees of image formation within the image-forming layers (i.e., different gray levels of the image) may be achieved by selecting a particular group of mini-subintervals, from among the total number of mini-subintervals available, during which a pulse of electrical current will be applied. The different degrees of image formation may be
20 achieved either by changing the size of dots printed in the image-forming layer(s), or by changing the optical density of dots printed in the image-forming layer(s), or by a combination of variations in dot size and optical density.

Although the method has been described above with reference to a single pixel, printed by a single heating element of the printhead, it will be apparent to one
25 of skill in the art that a printhead may contain a linear array of many such heating elements, and that the thermal imaging member may be translated beneath this linear array, in a direction orthogonal to said linear array, such that an image of a line of pixels may be formed in the thermal imaging member during the time interval for forming an image of a single pixel by a single heating element. Further, it will be
30 clear to one of skill in the art that images may be formed in either or both of the image-forming layers of the thermal imaging member during the time interval for

forming an image of a single pixel by a single heating element, the image in the first image-forming layer being formed by the energy applied during the first time interval specified above, and the image formed in the second image-forming layer being formed by the energy applied during the second time interval specified above.

5 Thus, both images may be formed when the thermal imaging member is translated once beneath the printhead, i.e., in a single pass of the printhead. In practice, the energy applied during the first time period will heat the second image-forming layer, and the energy applied during the second time period will heat the first image-forming layer. Those of skill in the art will appreciate that suitable adjustment of the energy supplied during both time periods will be required in order to compensate for
10 these effects, as well as to compensate for other effects, such as thermal history and unintended heating by adjacent heating elements.

In actual practice, the number of pulses can be quite different than that shown in Figs. 13 and 14. In a typical printing system, the pixel-printing interval
15 may be in the range of 1-100 milliseconds and the mini-subinterval length may be in the range of 1-100 microseconds. There are therefore typically hundreds of mini-subintervals within the pixel-printing interval.

The duty cycle within a mini-subinterval can generally be changed from pulse to pulse and, in another preferred embodiment, this technique may be used to
20 tailor the average power applied to the heating elements to achieve good printing results.

Of course, it will be apparent to those skilled in the art that where it is desired to address independently more than two image-forming layers of the imaging member in a single pass, the available number of mini-subintervals and the
25 range of duty cycles must be divided into a correspondingly larger number of combinations, each capable of printing at least partially independently on one of the image-forming layers.

In a particularly preferred embodiment of the invention, three different image-forming layers carried by the same surface of the substrate of the thermal
30 imaging member are addressed from the same surface of the imaging member by one thermal printhead in a single pass. This embodiment will be described in

relation to Fig. 9. The substrate 22 may be any of the materials previously described. Image-forming layer 28 comprises a meltable leuco dye having a melting point of from about 90°C to about 140°C and a developer material having a melting point in the same range, and optionally includes a thermal solvent having a melting point in the same range. In this embodiment layer 28 is about 1 to 4 μm thick and is coated from an aqueous dispersion. Interlayer 32 is about 5 to about 25 μm thick and comprises a water-soluble inert material which may be any suitable water-soluble interlayer material previously mentioned. The second image-forming layer, 26, comprises a leuco dye and a developer material, each having a melting point of from about 150°C to about 280°C, and optionally includes a thermal solvent having a melting point in the same range. The second image-forming layer has a thickness of from about 1 to about 4 μm and is coated from a water dispersion. The second interlayer, 30, comprises a water-soluble inert material, which may be any of the water-soluble interlayer materials previously mentioned, and has a thickness of from about 3 to about 10 μm. The third image-forming layer, 24, comprises either: a) a meltable leuco dye having a melting point of at least 150°C, preferably 250°C, and a developer material having a melting point of at least 250°C, preferably 300°C, optionally including a thermal solvent; or b) a molecule which forms color unimolecularly at a temperature of at least 300°C in about from 0.1 to about 2 milliseconds (a suitable material is Leuco Dye III described in detail below herein). The third image-forming layer has a thickness of from about 1 to about 4 μm and is coated from a water dispersion. This particularly preferred thermal imaging member further includes an overcoat layer such as is described in Example I below.

As described above, Figs 8 - 10 relate to a thermal imaging member for which thermal diffusion is the technique used for partitioning the time-temperature domain. Another technique for partitioning the time – temperature domains of a thermal imaging member in accordance with the invention resides in the exploitation of phase transitions. The phase transitions, for example, may be the result of a natural melting or glass transitions of the dye itself, or may be achieved by incorporating thermal solvents into the dye layers. When a measurement is made of

the time t required to reach a certain optical density of the dye when the dye layer is held at a fixed temperature T it is typically found that the relationship between the temperature and the time is expressed by an Arrhenius curve:

$$\log(t) \sim (-A + B/T)$$

5 where A and B are constants that may be determined experimentally. When measurements are taken in the temperature range of a melting transition, it is often found that the slope, B , far exceeds that normally found in regions removed from phase transitions. As a result, the Arrhenius curve for a normal dye layer (i.e., one in which no phase change is associated with imaging, as will be the case for
10 diffusion-controlled reactions, for example) and for a melting dye layer may cross at a steep angle, as shown in Fig. 15 for a cyan dye, namely 3-(1-n-butyl-2-methylindol-3-yl)-3-(4-dimethylamine-2-methylphenyl) phthalide, available from Hilton-Davis Company, in conjunction with a Lewis Acid developer, the zinc salt of 3,5-di-*t*-butylsalicylic acid and a naturally melting magenta dye, namely Solvent
15 Red 40, available from Yamamoto Chemical Company in conjunction with an acid developer, bis(3-allyl-4-hydroxyphenyl) sulfone, available from Nippon Kayaku Company, Ltd. The two curves show the time required to reach a density of 0.1 for each dye. Such a relationship may itself be used as the basis for a multicolor thermal printing system according to one embodiment of the present invention, insofar as Fig. 15 shows that below the crossing temperature the cyan dye turns on
20 more quickly than the magenta dye and above the crossing temperature the magenta dye turns on more quickly than the cyan dye. For the two dyes shown, it is seen that it would take more than one second per line to print cyan without magenta contamination. To overcome this limitation, the dyes or their environment may be
25 modified to move the crossing point to a shorter time region. However, the system may be made even more desirable from a time consideration by "burying" the magenta dye layer as described above in Fig 8.

Yet another technique for partitioning the time – temperature domains of a thermal imaging member in accordance with the invention is illustrated in Fig. 16.
30 This technique employs a multicolor thermal imaging member 60 according to the invention which includes a layer of a magenta image-forming material 62, in this

illustrative instance a leuco dye, associated with a layer 64 of an acid developer material having a melting point, T_7 and a layer of a cyan image forming material 66 associated with a layer 68 of an acid developer material having a melting point, T_8 . The imaging member 60 also includes first and second timing layers, 70 and 72, respectively, and a layer 74 of a fixing material having a melting point, T_9 . Imaging member 60 may also include a substrate (not shown) which may be positioned adjacent layer 64 or layer 68.

There are known leuco dyes that form color irreversibly upon contact with suitable developers. With this type of dye, layer 74 of fixing material functions to terminate, but not reverse, color formation in either of the two image-forming layers, 62 and 66, respectively. The fixing material, however, must pass through the timing layers, 70 and 72, respectively, by diffusion or dissolution to terminate color formation within the image-forming layers. As shown, one of the timing layers, in this illustrative instance timing layer 70, is thinner than the other timing layer 72 and therefore the fixing material arrives at cyan image-forming layer 66 later than when it arrives at magenta image-forming layer 62. Thus, a timing difference is introduced between the formation of the two colors in accordance with the invention.

The developer layers 64 and 68 must melt before the developer materials can combine with the leuco dyes. By selecting the materials in the developer layer such that they melt at different temperatures, a temperature difference is introduced between the formation of the two colors in accordance with the invention. In this illustrative embodiment T_7 is lower than T_8 , e.g., $T_7 = 120^\circ\text{C}$ and $T_8 = 140^\circ\text{C}$. In this embodiment of the invention various possibilities are provided. Where the imaging member is heated to a temperature less than 120°C , then neither of the developer layers, 64 and 68, will melt and no color will be formed. Further, provided that the thermal energy applied to the imaging member is sufficient to melt the fixing material, the melting point of the fixing layer, T_9 , being less than the melting points, T_7 and T_8 , respectively, of the developer layers, (e.g., $T_9 = 100^\circ\text{C}$) the fixing material will diffuse through the timing layers 70 and 72 and eventually fix both

image-forming layers so that subsequent temperature applications will not cause any color to form.

When the imaging member 60 is heated to a temperature between T_7 and T_8 then developer material in layer 64 will melt and begin to mix with the magenta leuco dye precursor to form color. The amount of color formation is dependent primarily upon the amount of time the temperature of the developer layer 64 remains above T_7 . Following this thermal exposure the temperature of the imaging member is lowered below T_7 and held at that temperature until the fixing material arrives and prevents any further color formation. When the temperature of the imaging member is held below T_7 for a longer period of time the fixing material will also arrive at the cyan image-forming layer 66 and prevent any future formation of color by this layer. In this manner a selectable amount of magenta color can be formed without forming any cyan color.

In a similar manner a selectable amount of cyan can be formed in accordance with the invention without forming any magenta. Initially, the imaging member is heated to a temperature above T_9 but below T_7 in order to allow the fixing material to arrive at magenta image-forming layer 62 and inactivate it, thereby preventing it from subsequently forming any color. Subsequently, the temperature is raised above T_8 to cause the developer material in layer 68 to combine with the leuco dye precursor and begin the formation of cyan color. The amount of cyan color formation is primarily dependent upon the amount of time the temperature of the imaging member is maintained above T_8 . It will be appreciated that this procedure will also cause the developer material in layer 64 to melt but no formation of magenta color results since the magenta dye precursor was previously fixed. Subsequently, the temperature of the imaging member 60 is lowered below T_7 and held at that level until the fixing material arrives at layer 66 to prevent the formation of any further cyan.

In order to print both magenta and cyan, the sequence of heat pulses applied to the imaging member 60 is such as to carry out a combination of the steps described above to create cyan and magenta, respectively. Initially, the imaging member 60 is heated to a temperature above T_7 to produce a selectable density of

magenta. The temperature is then lowered below T_7 for a period of time sufficient to fix the magenta precursor layer 62 followed by raising the temperature above T_8 to produce a selectable density of cyan color and then once again lowering the temperature below T_7 to fix the cyan precursor layer 66.

5 As previously described, a wide variety of different irreversible chemical reactions may be used to achieve a color change in a layer. The fixer material used in any particular instance will depend upon the choice of mechanism exploited to achieve the color change. For example, the mechanism may involve the coupling of two colorless materials to form a colored dye. In this case, the fixing reagent would
10 react with either of the two dye precursor molecules to form a colorless product thereby interfering with any further formation of dye.

 A negative working version of a two-color imaging member according to the invention may also be devised according to the same principles, as illustrated in Fig. 17. In this implementation the dye layers are initially colored, and they remain so
15 unless an adjacent layer of decolorizing reagent thermally activated before the arrival of the fixing reagent through a timing layer. Referring now to Fig. 17 there is seen a negative working thermal imaging member 80 according to the invention which includes a first image-forming layer 82, e.g., a magenta dye layer, a second image-forming layer 84, e.g., a cyan dye layer, first and second timing layers 86 and
20 88, respectively, a fixing layer 90 and first and second decolorizer layers 92 and 94, respectively. Imaging member 80 may also include a substrate (not shown) which may be positioned adjacent layer 92 or layer 94.

 For example, the magenta and cyan dyes may be irreversibly decolorized by exposure to a base as described in U.S. Patents Nos. 4,290,951 and 4,290,955.
25 Where the reagent layer 90 contains an acidic material and the acid is chosen so as to neutralize the basic material in the decolorizing layers 92 and 94, it will be appreciated that where the acid arrives in the dye-containing layers before the base, the base will not be able to decolorize the magenta or cyan dye whereas when the base arrives before the acid, irreversible decolorization will have occurred. As
30 discussed above in relation to the embodiment shown in Fig. 8, the third color may be obtained by any other printing modality including thermally printing the third

color from the back of the imaging member as described in relation to Figs. 9 and 10.

Fig. 18 illustrates a three-color thermal imaging member according to the invention. Referring now to Fig. 18 there is seen imaging member 100 which includes the layers shown for the imaging member 60 which is illustrated in Fig. 16 and these layers are designated by the same reference numerals. Imaging member 100 also includes a buffer layer 102, yellow dye precursor layer 104 and a third acid developer layer 106 in which the developer material has a melting point T_{10} which is higher than T_7 and T_8 . After forming the desired color densities in cyan and magenta as described above in relation to Fig. 16, the temperature of the imaging member can be raised above T_{10} to form a selectable density of yellow dye. It should be noted that where T_{10} is a temperature higher than the imaging member 100 is likely to encounter during its useful life, it is not necessary to inactivate the yellow dye precursor subsequent to writing the yellow image. Imaging member 100 may also include a substrate (not shown) which may be positioned adjacent layer 64 or layer 106.

In choosing the layer dimensions for the imaging members illustrated in Figs 16 and 18 it is advantageous to have the timing layer 70 be as thin as possible but not substantially thinner than dye layer 62. Timing layer 72 typically will be about two to three times the thickness of timing layer 70.

It will be appreciated that the practice of the invention according to the methods just described relies upon the diffusion or dissolution of chemical species, rather than the diffusion of heat. Whereas the thermal diffusion constant is normally relatively insensitive to temperature, the diffusion constants for chemical diffusion are typically exponentially dependent on the inverse of the temperature, and therefore more sensitive to changes in the ambient temperature. Moreover, when dissolution is chosen as the time-determining mechanism, numerical simulations show that the timing is typically quite critical because the colorization process occurs relatively quickly once the timing layer has been breached.

Any chemical reaction in which color is formed irreversibly is, in principle, amenable to the fixing mechanism described above. Materials that form color

irreversibly include those in which two materials couple together to form a dye. The fixing mechanism is achieved by introducing a third reagent that couples preferentially with one of the two dye-forming materials to form a colorless product.

5 In addition to the methods recited above, chemical thresholds can also be used to partition the time-temperature domain in accordance with the multicolor thermal imaging system of the invention. As an example of this mechanism, consider a leuco dye reaction in which the dye is activated when it is exposed to an acid. If, in addition to the dye, the medium contains a material significantly more basic than the dye, which does not change color when protonated by the acid,
10 addition of acid to the mixture will not result in any visible color change until all of the more basic material has been protonated. The basic material provides for a threshold amount of acid which must be exceeded before any coloration is evident. The addition of acid may be achieved by various techniques such as by having a dispersion of acid developer crystals which melt and diffuse at elevated temperatures
15 or by having a separate acid developer layer which diffuses or mixes with the dye layer when heated.

A certain time delay is involved in reaching the acid level required to activate the dye. This time period may be adjusted considerably by adding base to the imaging member. In the presence of added base, as described above, there is an
20 interval of time required for the increasing amount of acid to neutralize the base. Beyond this time period, the imaging member will be colorized. It will be seen that the same technique can be used in a reverse sequence. A dye that is activated by base can have its timing increased by the addition of a background level of acid.

In this particular embodiment, it is notable that the diffusion of the acid or
25 base developer material into the dye-containing layer is typically accompanied by diffusion of dye in reverse into the developer layer. When this occurs, color formation may begin almost immediately since the diffusing dye may find itself in an environment where the developer material level far exceeds the threshold level necessary to activate the dye. Accordingly, it is preferred to inhibit the dye from
30 diffusing into the developer layer. This may be accomplished, for example, by

attaching long molecular chains to the dyes, by attaching the dyes to a polymer, or by attaching the dye to an ionic anchor.

EXAMPLES

5 The thermal imaging system of the invention will now be described further with respect to specific preferred embodiments by way of examples, it being understood that these are intended to be illustrative only and the invention is not limited to the materials, amounts, procedures and process parameters, etc. recited therein. All parts and percentages are by weight unless otherwise specified.

The following materials were used in the examples described below:

10 Leuco Dye I, 3,3-bis(1-n-butyl-2-methyl-indol-3-yl)phthalide (Red 40, available from Yamamoto Chemical Industry Co., Ltd., Wakayama, Japan);

Leuco Dye II, 7-(1-butyl-2-methyl-1H-indol-3-yl)-7-(4-diethylamino-2-methyl-phenyl)-7H-furo[3,4-b]pyridin-5-one (available from Hilton-Davis Co., Cincinnati, OH);

15 Leuco Dye III, 1-(2,4-dichloro-phenylcarbamoyl)-3,3-dimethyl-2-oxo-1-phenoxy-butyl]-(4-diethylamino-phenyl)-carbamic acid isobutyl ester, prepared as described in U. S. Patent No. 5,350,870;

Leuco Dye IV, Pergascript Yellow I-3R, available from Ciba Specialty Chemicals Corporation, Tarrytown, NY;

20 Acid Developer I, bis(3-allyl-4-hydroxyphenyl)sulfone, available from Nippon Kayaku Co., Ltd, Tokyo, Japan;

Acid Developer II, PHS-E, a grade of poly(hydroxy styrene), available from TriQuest, LP, a subsidiary of ChemFirst Inc., Jackson, MS;

25 Acid Developer III, zinc salt of 3,5-di-t-butyl salicylic acid, available from Aldrich Chemical Co., Milwaukee, WI;

Acid Developer IV, zinc salt of 3-octyl-5-methyl salicylic acid, prepared as described in Example 7 below;

Airvol 205, a grade of poly(vinyl alcohol) available from Air Products and Chemicals, Inc., Allentown, PA;

Airvol 350, a grade of poly(vinyl alcohol) available from Air Products and Chemicals, Inc., Allentown, PA;

Airvol 540, a grade of poly(vinyl alcohol) available from Air Products and Chemicals, Inc., Allentown, PA;

5 Genflo 305, a latex binder, available from Omnova Solutions, Fairlawn, OH;
 Genflo 3056, a latex binder, available from Omnova Solutions, Fairlawn,
 OH;

 Glascol C44, an aqueous polymer dispersion, available from Ciba Specialty Chemicals Corporation, Tarrytown, NY;

10 Joncryl 138, a binder, available from S. C. Johnson, Racine, WI;

 Irganox 1035, an antioxidant, available from Ciba Specialty Chemicals Corporation, Tarrytown, NY;

 Aerosol-OT, a surfactant available from Dow Chemical, Midland, MI;

 Dowfax 2A1, a surfactant available from Dow Chemical Corporation,
15 Midland, MI;

 Ludox HS40, a colloidal silica available from DuPont Corporation,
 Wilmington, DE;

 Nipa Proxel, a bactericide available from Nipa Inc., Wilmington, DE;

 Pluronic 25R2, a surfactant available from BASF, Ludwigshaven, Germany;

20 Tamol 731, a polymeric surfactant (sodium salt of polymeric carboxylic acid) available from Rohm and Haas Company, Philadelphia, PA;

 Triton X-100, a surfactant available from Dow Chemical Corporation,
 Midland, MI;

 Zonyl FSN, a surfactant, available from DuPont Corporation, Wilmington,
25 DE;

 Zonyl FSA, a surfactant, available from DuPont Corporation, Wilmington,
 DE;

 Hymicron ZK-349, a grade of zinc stearate available from Cytech Products, Inc., Elizabethtown, KY;

30 Klebosol 30V-25, a silica dispersion available from Clariant Corporation,
 Muttens, Switzerland;

Titanium dioxide, a pigment available from DuPont Corporation,
Wilmington, DE;

Glyoxal, available from Aldrich Chemical Co., Milwaukee, WI;

Melinex 534, a white poly(ethylene terephthalate) film base of
5 approximately 96 microns' thickness, available from DuPont Corporation,
Wilmington, DE);

Cronar 412, a clear poly(ethylene terephthalate) film base of approximately
102 microns' thickness, available from DuPont Corporation, Wilmington, DE.

EXAMPLE I

10 A two color imaging member such as is illustrated in Fig. 8 and further
including an overcoat layer deposited on the cyan color-forming layer was prepared
as follows:

A. The magenta image-forming layer was prepared as follows:

A leuco magenta dye, Leuco Dye I, was dispersed in an aqueous mixture
15 comprising Airvol 205 (4.5% of total solids) and surfactants Pluronic 25R2 (1.5%
of total solids) and Aerosol-OT (5.0% of total solids) in deionized water, using an
attriter equipped with glass beads, stirred for 18 hours at 2 °C. The average particle
size of the resulting dispersion was about 0.28 microns and the total solid content
was 19.12%.

20 Acid Developer I was dispersed in an aqueous mixture comprising Airvol
205 (7.0% of total solids), Pluronic 25R2 (1.5% of total solids), and deionized water,
using an attriter equipped with glass beads and stirred for 18 hours at 2°C. The
average particle size of the resulting dispersion was about 0.42 microns, and the total
solid content was 29.27%.

25 The above dispersions were used to make the magenta coating fluid in
proportions stated below. The coating composition thus prepared was coated onto
Melinex 534 using a Meyer rod, and dried. The intended coating thickness was 2.9
microns.

Ingredient	% solids in dried film
Leuco Dye I	10.74%
Acid Developer I	42.00%
Genflo 3056	47.05%
Zonyl FSN	0.21%

B. A thermally insulating interlayer was deposited onto the magenta imaging layer as follows:

A coating fluid for the interlayer was prepared in proportions stated below. The image interlayer coating composition thus prepared was coated on the magenta imaging layer using a Meyer rod for an intended thickness of 13.4 microns, and was dried in air.

Ingredient	% solids in dried film
Glascot C44	99.50%
Zonyl FSA	0.50%

C. Cyan image-forming layers C1 – C3 were deposited on the thermally insulating layer as follows:

C1 Cyan developer layer.

10 Acid Developer III was dispersed in an aqueous mixture comprising of Airvol 205 (6.0% of total solids), Aerosol-OT (4.5% of total solids) and Triton X-100 (0.5% of total solids) in deionized water, using an attriter equipped with glass beads, by stirring for 18 hours at room temperature. The average particle size of the resulting dispersion was about 0.24 microns, and the total solid content was 25.22%.

15 The above dispersion was used to make the cyan developer coating fluid in proportions stated below. The cyan developer coating composition thus prepared was coated on top of the imaging interlayer using a Meyer rod for an intended thickness of 1.9 microns, and was dried in air.

Ingredient	% solids in dried film
Joncyl 138	9.50%
Acid Developer III	89.50%
Zonyl FSN	1.00%

C2 Cyan interlayer.

A cyan interlayer coating fluid was prepared in proportions stated below. The cyan interlayer coating composition thus prepared was coated on top of the cyan developer layer using a Meyer rod for an intended thickness of 2.0 microns, and was
5 dried in air.

Ingredient	% solids in dried film
Airvol 205	99.00%
Zonyl FSN	1.00%

C3 Cyan dye layer.

The leuco cyan dye, Leuco Dye II, was dispersed in an aqueous mixture comprising Airvol 350 (7.0% of total solids), Airvol 205 (3.0% of total solids), Aerosol-OT (1.0% of total solids) and Triton X-100 (0.2% of total solids) in
10 deionized water, using an attriter equipped with glass beads, stirred for 18 hours at room temperature. The average particle size of the resulting dispersion was about 0.58 microns, and the total solid content was 26.17%.

The above dispersion was used to make the cyan coating fluid in proportions stated below. The cyan coating composition thus prepared was coated on the cyan
15 interlayer using a Meyer rod for an intended thickness of 0.6 microns, and was dried in air.

Ingredient	% solids in dried film
Leuco Dye II	59.5%
Joncryl 138	39.5%
Zonyl FSN	1.0%

D. A protective overcoat was deposited on the cyan color-forming layers as follows:

A slip overcoat was coated on the cyan dye layer. The overcoat was
20 prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the cyan dye layer using a Meyer rod for an intended thickness of 1.0 micron, and was dried in air.

Ingredient	% solids in dried film
Glyoxal	9.59%
Hymicron ZK-349	31.42%
Klebosol 30V-25	23.53%
Zonyl FSA	3.89%
Airvol 540	31.57%

The resulting six-layer imaging member was printed using a laboratory test-bed printer equipped with a thermal head, model KST-87-12MPC8 (Kyocera Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan).

The following printing parameters were used:

- 5 Printhead width: 3.41 inch
- Pixels per inch: 300
- Resistor size: 69.7 x 80 microns
- Resistance: 3536 Ohm
- Line Speed: 8 milliseconds per line
- 10 Print speed: 0.42 inches per second
- Pressure: 1.5 - 2 lb/linear inch
- Dot pattern: Rectangular grid.

The cyan layer was printed with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.3 milliseconds (about 16.3% of the total line time) in twenty equal steps, while the voltage supplied to the print head was maintained at 27.0V.

A lower power/longer time condition was used to print the magenta layer. The pulse width was increased from zero to the full 8 millisecond line time in twenty equal steps, while the voltage supplied to the print head was maintained at 14.5V.

20 Following printing, the reflection density in each of the printed areas was measured using a spectrophotometer from GretagMacbeth AG, Regensdorf, Switzerland. The results are shown in Tables I and II. Table I shows the printing of the cyan layer as a function of energy supplied by the thermal head. The magenta densities obtained are shown as well. Also included in Table I is the ratio between the cyan and the magenta density (C/M). Similarly, Table II shows the printing of

the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the cyan densities is shown (M/C).

The ratio C/M in Table I and the ratio M/C in Table II are measured quantities that indicate success in differentially printing one color rather than another. However, there are two reasons why these numbers do not fully reflect the degree of layer discrimination. First, the measured densities have a contribution resulting from absorption of light by the underlying media substrate. (For example, even in the absence of printing there is a residual absorption of 0.04 density units.) Second, each of the dyes has some absorption outside of its own color band. Therefore, the ratio of measured cyan and magenta optical densities is not the same as the ratio of colorized cyan dye to colorized magenta dye.

An approximate correction for substrate absorption may be made by subtracting the optical density of the unheated media from each of the measured density values. Correcting for the out-of-band absorption of each of the dyes is more complicated. Here there is considered a three-color imaging member (comprised of three dye layers) as a general example for the correction procedure,

First, the out-of-band absorption was characterized by measuring the density of each of the three dyes in each of the three color bands, and correcting the densities for the substrate density. Three monochrome samples were used, and each had a particular area-concentration a_j^0 of one of the dyes, where $j = C, M$ or Y depending on whether the dye was cyan, magenta or yellow, respectively.

The results of such a measurement were:

	Cyan Dye	Magenta Dye	Yellow Dye
Cyan Density	0.75	0.02	0.00
Magenta Density	0.26	0.63	0.04
Yellow Density	0.14	0.11	0.38

The densities recorded in this matrix will be denoted d_{ij} , where i and j are the color values C, M and Y, and for example the value d_{CM} is the magenta density of the cyan dye sample

5 If we have colorized dyes of area-concentration other than that at which these data were recorded, then the densities for that dye will scale in proportion to the area-concentration. In particular, if a sample has area concentrations a_C , a_M , and a_Y of colorized cyan, magenta and yellow dye, then under the same printing conditions we will observe measured densities D_C , D_M and D_Y of

$$\begin{aligned} 10 \quad D_C &= (a_C/a_C^0) d_{CC} + (a_M/a_M^0) d_{MC} + (a_Y/a_Y^0) d_{YC} \\ D_M &= (a_C/a_C^0) d_{CM} + (a_M/a_M^0) d_{MM} + (a_Y/a_Y^0) d_{YM} \\ D_Y &= (a_C/a_C^0) d_{CY} + (a_M/a_M^0) d_{MY} + (a_Y/a_Y^0) d_{YY} \end{aligned}$$

This can be written in standard matrix notation in the following way:

$$15 \quad \begin{pmatrix} D_C \\ D_M \\ D_Y \end{pmatrix} = \begin{pmatrix} d_{CC} & d_{MC} & d_{YC} \\ d_{CM} & d_{MM} & d_{YM} \\ d_{CY} & d_{MY} & d_{YY} \end{pmatrix} \begin{pmatrix} a_C/a_C^0 \\ a_M/a_M^0 \\ a_Y/a_Y^0 \end{pmatrix}$$

If the densities D_C , D_M and D_Y of a sample are measured, then we can use the inverse of this equation to find the area concentrations of colorized dye in the sample, in comparison to those of the calibration samples.

$$\begin{pmatrix} a_C/a_C^0 \\ a_M/a_M^0 \\ a_Y/a_Y^0 \end{pmatrix} = \begin{pmatrix} d_{CC} & d_{MC} & d_{YC} \\ d_{CM} & d_{MM} & d_{YM} \\ d_{CY} & d_{MY} & d_{YY} \end{pmatrix}^{-1} \begin{pmatrix} D_C \\ D_M \\ D_Y \end{pmatrix}$$

25 These quantities more accurately represent the colorization of each layer by the applied heat, and are not confounded by the spectral absorption overlaps of the dyes in those layers. As such, they more accurately represent the degree to which we are able to write on one layer without affecting another.

We can define “cross-talk” to be the degree to which an attempt to produce optical density in one color layer alone results in the production of undesired optical

density in another color layer. For example, if we have a medium with a cyan layer and a magenta layer, and we are attempting to write on the magenta layer, then the relative cross-talk from cyan may be represented by:

$$5 \quad \text{Cross - talk} = \frac{a_C * (d_{CC} / a_C^0)}{a_M * (d_{MM} / a_M^0)} = \frac{a_C / a_C^0}{a_M / a_M^0} \left(\frac{d_{CC}}{d_{MM}} \right)$$

An analogous equation can be written for the cross-talk of magenta when attempting to write on the cyan layer.

- These values of cross-talk are recorded in the final column of Tables I and II.
- 10 Similar values will be reported for the following examples as well, but only for cases in which the measured densities are large enough (density > 0.1) to yield meaningful results, and only for layers that are addressed from the same surface of the imaging member.

Table I

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	C/M	Cross-Talk (Magenta)
0.00	0.04	0.04	1.00	
0.18	0.04	0.04	1.00	
0.35	0.04	0.04	1.00	
0.53	0.04	0.04	1.00	
0.71	0.04	0.04	1.00	
0.88	0.04	0.04	1.00	
1.06	0.04	0.04	1.00	
1.24	0.04	0.04	1.00	
1.41	0.04	0.05	0.80	
1.59	0.05	0.05	1.00	
1.77	0.06	0.05	1.20	
1.94	0.1	0.06	1.67	
2.12	0.15	0.08	1.88	
2.29	0.2	0.1	2.00	
2.47	0.29	0.12	2.42	0.01
2.65	0.34	0.15	2.27	0.04
2.82	0.43	0.22	1.95	0.14
3.00	0.5	0.29	1.72	0.22
3.18	0.62	0.35	1.77	0.22
3.35	0.6	0.42	1.43	0.37
3.53	0.61	0.47	1.30	0.45

Table II

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	M/C	Cross-Talk (Cyan)
0	0.04	0.04	1.00	
0.30	0.04	0.04	1.00	
0.60	0.04	0.05	1.25	
0.90	0.04	0.05	1.25	
1.21	0.04	0.05	1.25	
1.51	0.04	0.05	1.25	
1.81	0.04	0.05	1.25	
2.11	0.04	0.05	1.25	
2.41	0.05	0.06	1.20	
2.71	0.05	0.1	2.00	0.14
3.02	0.05	0.15	3.00	0.07
3.32	0.06	0.22	3.67	0.08
3.62	0.07	0.29	4.15	0.09
3.92	0.09	0.42	4.67	0.10
4.22	0.1	0.54	5.40	0.09
4.52	0.13	0.69	5.31	0.11
4.83	0.16	0.97	6.06	0.10
5.13	0.22	1.32	6.00	0.11
5.43	0.26	1.56	6.00	0.12
5.73	0.31	1.69	5.45	0.14
6.03	0.34	1.74	5.12	0.15

EXAMPLE II

5 This example illustrates a two-color imaging member such as is illustrated in Fig. 8. The top color-forming layer produces a yellow color, using a unimolecular thermal reaction mechanism as described in U. S. Patent No. 5,350,870. The lower color-forming layer produces a magenta color, using an acid developer and a magenta leuco dye.

10 A. The magenta image-forming layer was prepared as follows:

Dispersions of Leuco Dye I and Acid Developer I were prepared as described in Example I, part A above.

Acid Developer II was dispersed in an aqueous mixture comprising Airvol 205 (2% of total solids), Dowfax 2A1 (2% of total solids) and Irganox 1035 (5% of total solids) in deionized water, using an attriter equipped with glass beads and stirred for 24 hours at 10-15 °C. The average particle size of the resulting dispersion was about 0.52 microns and the total solid content was 22.51%.

The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated onto Melinex 534 using a Meyer rod, and dried. The intended coating thickness was 3 microns.

Ingredient	% solids in dried film
Leuco Dye I	24.18%
Acid Developer I	47.49%
Acid Developer II	11.63%
Joncryl 138	16.16%
Zonyl FSN	0.54%

B. A thermally insulating interlayer was deposited onto the magenta imaging layer as described in Example I, part B. above, except that the coating thickness was 16.1 microns.

C. A yellow image-forming layer was deposited on the thermally insulating layer as follows:

Leuco Dye III was dispersed in an aqueous mixture comprising of Airvol 205 (4.54% of total solids), Aerosol-OT (2.73% of total solids) and Pluronic 25R2 (1.82% of total solids) in deionized water, using an attriter equipped with glass beads and stirred for 18 hours at room temperature. The average particle size of the resulting dispersion was about 0.49 microns and the total solid content was 25.1%.

The above dispersion was used to make the yellow coating fluid in proportions stated below. The yellow coating composition thus prepared was coated on the thermally insulating interlayer using a Meyer rod for an intended thickness of 3 microns, and was dried in air.

Ingredient	% solids in dried film
Leuco Dye III	70%
Genflo 3056	22.95%
Airvol 205	7%
Zonyl FSN	0.05%

D. A protective overcoat was deposited on the yellow color-forming layer as follows:

A slip overcoat was coated on the yellow dye layer. The overcoat was prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the yellow dye layer using a Meyer rod for an intended thickness of 1.0 micron, and was dried in air.

Ingredient	% solids in dried film
Glyoxal	8.39%
Hymicron ZK-349	31.77%
Kiebosol 30R 25	23.77%
Zonyl FSA	0.92%
Zonyl FSN	3.22%
Airvol 540	31.93%

The resulting four-layer imaging member was printed using a laboratory test-bed printer equipped with a thermal head, model KST-87-12MPC8 (Kyocera Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan). The following printing parameters were used:

Printhead width: 3.41 inch
 Pixels per inch: 300
 Resistor size: 69.7 x 80 microns
 Resistance: 3536 Ohm
 Line Speed: 8 milliseconds per line
 Print speed: 0.42 inches per second
 Pressure: 1.5 - 2 lb/linear inch
 Dot pattern: Rectangular grid.

The yellow layer was printed with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a

maximum of 1.65 milliseconds (about 20.6% of the total line time) in twenty-one equal steps, while the voltage supplied to the print head was maintained at 29.0V.

A lower power/longer time condition was used to print the magenta layer. The pulse width was increased from zero to the 99.5% of the 8 millisecond line time
5 in twenty-one equal steps, while the voltage supplied to the print head was maintained at 16V.

Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. The results are shown in Tables III and IV. Table III shows the printing of the yellow layer as a function of
10 energy supplied by the thermal head. The magenta densities obtained are shown as well. Also included in Table III are the ratio between the yellow and the magenta density (Y/M) and the cross-talk.. Similarly, Table IV shows the printing of the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the yellow densities is shown (M/Y) as well as the cross-
15 talk..

Table III

Energy Supplied (J/cm ²)	Yellow printed density	Magenta printed density	Y/M	Cross-Talk (Magenta)
0.00	0.07	0.09	0.78	
0.26	0.07	0.09	0.78	
0.52	0.06	0.09	0.67	
0.78	0.06	0.09	0.67	
1.04	0.06	0.09	0.67	
1.30	0.07	0.09	0.78	
1.56	0.06	0.09	0.67	
1.82	0.06	0.09	0.67	
2.08	0.08	0.09	0.89	
2.34	0.11	0.10	1.10	
2.60	0.17	0.10	1.70	
2.86	0.24	0.11	2.18	0.01
3.12	0.34	0.12	2.83	0.01
3.38	0.48	0.14	3.43	0.02
3.64	0.58	0.16	3.63	0.03
3.90	0.68	0.19	3.58	0.06
4.16	0.83	0.23	3.61	0.08
4.41	0.94	0.26	3.62	0.09
4.67	1.08	0.32	3.38	0.13
4.93	1.13	0.38	2.97	0.18
5.19	1.19	0.40	2.98	0.18

Table IV

Energy Supplied (J/cm ²)	Magenta printed density	Yellow printed density	M/Y	Cross-Talk (Yellow)
0.00	0.10	0.08	1.25	
0.38	0.10	0.09	1.11	
0.76	0.10	0.09	1.11	
1.15	0.10	0.09	1.11	
1.53	0.10	0.08	1.25	
1.91	0.10	0.08	1.25	
2.29	0.10	0.07	1.43	
2.67	0.10	0.07	1.43	
3.05	0.10	0.07	1.43	
3.44	0.10	0.09	1.11	
3.82	0.10	0.08	1.25	
4.20	0.11	0.08	1.38	
4.58	0.14	0.1	1.40	
4.96	0.23	0.13	1.77	
5.35	0.40	0.18	2.22	0.22
5.73	0.61	0.25	2.44	0.17
6.11	0.88	0.34	2.59	0.17
6.49	1.17	0.44	2.66	0.17
6.87	1.42	0.53	2.68	0.17
7.26	1.65	0.65	2.54	0.20
7.64	1.68	0.74	2.27	0.26

EXAMPLE III

5 This example illustrates a two-color imaging member such as is illustrated in Fig. 8 and further including an overcoat layer deposited on the cyan color-forming layer. In this example, the thermally-insulating layer 18 of Fig. 8 is opaque, while the substrate 12 is transparent. It is therefore possible, using the imaging member described in this example, to print both sides of an opaque imaging member

10 independently, using a thermal head located on only one side of the imaging member.

A. Dispersions of Leuco Dye I and Acid Developer I were prepared as described in Example IV, part C below.

 Acid Developer II was dispersed as described above in Example II, part A.

The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated onto clear polyester film base (Cronar 412), and dried. The intended coating coverage was 3.3 g/m².

Ingredient	% solids in dried film
Leuco Dye I	21.91%
Acid Developer I	52.71%
Airvol 205	14.35%
Acid Developer II	10.54%
Zonyl FSN	0.49%

- 5 B. A thermally insulating interlayer was deposited onto the magenta imaging layer as follows:

A coating fluid for the interlayer was prepared in proportions stated below. The image interlayer coating composition thus prepared was coated on the magenta imaging layer for an intended thickness of 8.95 microns.

Ingredient	% solids in dried film
Glascol C44	99.50%
Zonyl FSA	0.50%

- 10 C. An opaque layer was deposited onto the thermally-insulating layer as follows:

A dispersion of titanium dioxide was prepared as follows:

- 15 Titanium dioxide was dispersed in an aqueous mixture comprising Tamol 731 (3.86% of total solids), Ludox HS40 (3.85% of total solids) and a trace amount (750ppm) of Nipa Proxel in deionized water, using an attriter equipped with glass beads and stirred for 18 hours at room temperature. The total solid content of the dispersion was 50.2%.

- 20 The dispersion so prepared was used to make a coating fluid in the proportions shown below. The coating fluid was coated onto the thermally-insulating layer for an intended thickness of 12.4 microns.

Ingredient	% solids in dried film
Titanium Dioxide	81.37%
Joncryl 138	18.08%
Zonyl FSN	0.54%

D. Cyan image-forming layers D1 – D3 were deposited on the thermally insulating layer as follows:

D1 Cyan developer layer.

5 Acid Developer III was dispersed as described in Example IV, part E1 below.

The above dispersion was used to make the cyan developer coating fluid in proportions stated below. The cyan developer coating composition thus prepared was coated on top of the imaging interlayer for an intended thickness of 1.74
10 microns.

Ingredient	% solids in dried film
Acid Developer III	80.84%
Joncryl 138	18.54%
Zonyl FSN	0.62%

D2 Cyan interlayer.

A cyan interlayer coating fluid was prepared in proportions stated below. The cyan interlayer coating composition thus prepared was coated on top of the cyan developer layer for an intended thickness of 1.0 microns.

Ingredient	% solids in dried film
Airvol 205	99.00%
Zonyl FSN	1.00%

15 D3 Cyan dye layer.

The leuco cyan dye, Dye II, was dispersed as described in Example 4, part E3 below.

The dispersion was used to make the cyan coating fluid in proportions stated below. The cyan coating composition thus prepared was coated on the cyan
20 interlayer for an intended thickness of 0.65 microns.

Ingredient	% solids in dried film
Dye II	59.30%
Joncryl 138	39.37%
Zonyl FSN	1.33%

E. A protective overcoat was deposited on the cyan color-forming layers as follows:

A slip overcoat was coated on the cyan dye layer. The overcoat was prepared in proportions stated in Table VI. The overcoat coating composition thus prepared was coated on the cyan dye layer for an intended thickness of 1.1 micron.

Ingredient	% solids in dried film
Hymicron ZK-349	31.77%
Klebosol 30V-25	23.77%
Airvol 540	31.93%
Glyoxal	8.39%
Zonyl FSA	0.92%
Zonyl FSN	3.22%

The resulting imaging member was printed as described in Example II above. The cyan image was visible from the front of the substrate, while the magenta image was visible from the rear. Therefore, optical densities for the cyan image were obtained from the top surface of the imaging member, and optical densities for the magenta image from the rear of the imaging member.

The cyan layer was printed with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.41 milliseconds (about 18.5% of the total line time) in twenty equal steps, while the voltage supplied to the print head was maintained at 29.0V.

A lower power/longer time condition was used to print the magenta layer. The pulse width was increased from zero to the full 8 millisecond line time in twenty equal steps, while the voltage supplied to the print head was maintained at 14.5V.

Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. The results are shown in Tables V and VI. Table V shows the printing of the cyan layer as a function of energy supplied by the thermal head. The magenta densities obtained are shown as

well. Also included in Table V are the ratio between the cyan and the magenta density (C/M) and the cross-talk. Similarly, Table VI shows the printing of the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the cyan densities is shown (M/C), as well as the cross-talk.

Table V

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	C/M	Cross-Talk (Magenta)
0.00	0.08	0.08	1.00	
0.23	0.08	0.08	1.00	
0.47	0.08	0.08	1.00	
0.70	0.08	0.08	1.00	
0.93	0.08	0.08	1.00	
1.17	0.08	0.08	1.00	
1.40	0.08	0.08	1.00	
1.64	0.08	0.08	1.00	
1.87	0.08	0.09	0.89	
2.10	0.08	0.08	1.00	
2.34	0.09	0.09	1.00	
2.57	0.09	0.09	1.00	
2.80	0.1	0.09	1.11	
3.04	0.11	0.10	1.10	
3.27	0.13	0.10	1.30	
3.51	0.22	0.13	1.69	0.03
3.74	0.27	0.15	1.80	0.04
3.97	0.35	0.18	1.94	0.04
4.21	0.36	0.20	1.80	0.10
4.44	0.42	0.24	1.75	0.15
4.67	0.51	0.28	1.82	0.14

Table VI

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	M/C	Cross-Talk (Cyan)
0.00	0.08	0.11	1.38	
0.31	0.08	0.11	1.38	
0.63	0.08	0.11	1.38	
0.94	0.08	0.11	1.38	
1.25	0.08	0.11	1.38	
1.57	0.08	0.11	1.38	
1.88	0.08	0.11	1.38	
2.20	0.08	0.11	1.38	
2.51	0.08	0.11	1.38	
2.82	0.08	0.11	1.38	
3.14	0.08	0.11	1.38	
3.45	0.08	0.11	1.38	
3.76	0.08	0.11	1.38	
4.08	0.08	0.12	1.50	
4.39	0.09	0.12	1.33	
4.70	0.09	0.13	1.44	
5.02	0.10	0.18	1.80	0.27
5.33	0.12	0.25	2.08	0.27
5.65	0.13	0.36	2.77	0.18
5.96	0.16	0.59	3.69	0.14
6.27	0.19	0.76	4.00	0.14

EXAMPLE IV

5 A three-color imaging member such as is illustrated in Fig. 9 and further including an overcoat layer deposited on the cyan color-forming layer was prepared as follows:

A. A yellow image-forming layer was prepared as follows:

10 A leuco yellow dye, Leuco Dye IV, was dispersed by a method analogous to that used to provide the dispersion of Leuco Dye I in part C., below, to give a dye concentration of 20.0%.

 Acid Developer IV (10 g) was dispersed in an aqueous mixture comprising Tamol 731 (7.08 g of a 7.06% aqueous solution) and deionized water, 32.92 grams,

in a 4 ounce glass jar containing 10 grams Mullite beads, stirred for 16 hours at room temperature. The developer concentration was 20.0%.

The above dispersions were used to make the yellow coating fluid in proportions stated below. The coating composition thus prepared was coated onto Melinex 534, and dried. The intended coating coverage was 2.0 g/m².

Ingredient	% solids in dried film
Leuco Dye IV	41.44%
Acid Developer IV	41.44%
Joncryn 138	16.57%
Zonyl FSN	0.55%

B. A thermally insulating interlayer was deposited onto the yellow imaging layer as follows:

A coating fluid for the interlayer was prepared in proportions stated in Table II. The image interlayer coating composition thus prepared was coated on the yellow imaging layer for an intended coverage of 9.0 g/m².

Ingredient	% solids in dried film
Glascot C44	99.50%
Zonyl FSA	0.50%

C. The magenta image-forming layer was prepared as follows:

Leuco Dye I (15.0 g) was dispersed in an aqueous mixture comprising Airvol 205 (3.38 g of a 20% aqueous solution), Triton X-100 (0.6 g of a 5% aqueous solution), and Aerosol-OT (15.01 g of a 19% aqueous solution) in deionized water (31.07 g), in a 4 ounce glass jar containing Mullite beads, stirred for 16 hours at room temperature. The total dye content was 20.00%.

Acid developer I (10 g) was dispersed in an aqueous mixture comprising Tamol 731 (7.08 g of a 7.06% aqueous solution) and deionized water, 32.92 grams, in a 4 ounce glass jar containing 10 grams Mullite beads, stirred for 16 hours at room temperature. The developer concentration was 20.0%.

Acid developer II was dispersed as described above in Example II, part A.

The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated onto

the thermally-insulating interlayer, and dried. The intended coating coverage was 1.67 g/m².

Ingredient	% solids in dried film
Leuco Dye I	24.18%
Acid Developer I	47.50%
Joncryl 138	16.16%
Acid Developer II	11.63%
Zonyl FSN	0.54%

D. A thermally insulating interlayer was deposited onto the magenta imaging layer as follows:

- 5 A coating fluid for the interlayer was prepared in proportions stated below. The image interlayer coating composition thus prepared was coated on the magenta imaging layer in three passes, for an intended coverage of 13.4 g/m².

Ingredient	% solids in dried film
Glascol C44	99.50%
Zonyl FSA	0.50%

E. Cyan image-forming layers E1 – E3 were deposited on the thermally-insulating layer as follows:

- 10 E1 Cyan developer layer.

Acid developer III (10 g) was dispersed in an aqueous mixture comprising Tamol 731 (7.08 g of a 7.06% aqueous solution) and deionized water, 32.92 grams, in a 4 ounce glass jar containing 10 grams Mullite beads, stirred for 16 hours at room temperature. The developer concentration was 20.0%.

- 15 The above dispersion was used to make the cyan developer coating fluid in proportions stated below. The cyan developer coating composition thus prepared was coated on top of the thermally-insulating interlayer for an intended thickness of 1.94 g/m².

Ingredient	% solids in dried film
Acid Developer III	89.5%
Joncryl 138	9.5%
Zonyl FSN	1.0%

E2 Cyan interlayer.

A cyan interlayer coating fluid was prepared in proportions stated below. The cyan interlayer coating composition thus prepared was coated on top of the cyan developer layer for an intended thickness of 1.0 g/m².

Ingredient	% solids in dried film
Airvol 205	99.00%
Zonyl FSN	1.00%

5 E3 Cyan dye layer.

Leuco Dye II (15.0 g) was dispersed in an aqueous mixture comprising Airvol 350 (11.06 g of a 9.5% aqueous solution), Airvol 205 (2.25 g of a 20% aqueous solution), Aerosol-OT (2.53 g of a 19% aqueous solution) and Triton X-100 (1.49 g of a 5% aqueous solution) in deionized water (52.61 g) in a 4 ounce glass jar containing Mullite beads, stirred for 16 hours at room temperature. The dye concentration was 20.0%.

The above dispersion was used to make the cyan coating fluid in proportions stated below. The cyan coating composition thus prepared was coated on the cyan interlayer for an intended coverage of 0.65 g/m².

Ingredient	% solids in dried film
Leuco Dye II	59.30%
Joncryn 138	39.37%
Zonyl FSN	1.33%

15 F. A protective overcoat was deposited on the cyan color-forming layers as follows:

A slip overcoat was coated on the cyan dye layer. The overcoat was prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the cyan dye layer for an intended coverage of 1.1 g/m².

Ingredient	% solids in dried film
Hymicron ZK-349	31.77%
Klebosol 30V-25	23.77%
Airvol 540	31.93%
Glyoxal	8.39%
Zonyl FSA	0.92%
Zonyl FSN	3.22%

The resulting imaging member was printed using a laboratory test-bed printer equipped with a thermal head, model KST-87-12MPC8 (Kyocera Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan). The following printing parameters were used:

- 5 Printhead width: 3.41 inch
- Pixels per inch: 300
- Resistor size: 69.7 x 80 microns
- Resistance: 3536 Ohm
- Line Speed: 8 milliseconds per line
- 10 Print speed: 0.42 inches per second
- Pressure: 1.5 - 2 lb/linear inch
- Dot pattern: Rectangular grid.

The cyan layer was printed with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.31 milliseconds (about 16.4% of the total line time) in ten equal steps, while the voltage supplied to the print head was maintained at 29.0V.

A lower power/longer time condition was used to print the magenta layer. The pulse width was increased from zero to the 99.5% of the 8 millisecond line time in ten equal steps, while the voltage supplied to the print head was maintained at 15V.

A very low power/very long time was used to print the yellow layer. Some of the printing conditions were changed, as follows:

- Line Speed: 15.23 milliseconds per line
- Pulse width: 15.23 milliseconds
- 25 Print speed: 0.0011 inches per second

Lines printed: 1600, one step of maximum density.

Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. The results are shown in Tables VII, VIII and IX. Table VII shows the printing of the cyan layer as a function of energy supplied by the thermal head. The magenta and yellow densities and cross-talk obtained are shown as well. Similarly, Table VIII shows the printing of the magenta layer as a function of the energy supplied by the thermal head. Table IX shows the density obtained when printing the yellow layer as a function of applied voltage and energy.

10

Table VII

	Cyan printed density	Magenta printed density	Yellow printed density	Cross-Talk (Magenta)	Cross-Talk (Yellow)
0.00	0.06	0.07	0.17		
0.41	0.06	0.07	0.17		
0.83	0.06	0.07	0.17		
1.24	0.05	0.07	0.16		
1.65	0.06	0.07	0.16		
2.07	0.06	0.07	0.18		
2.48	0.07	0.08	0.19		
2.89	0.12	0.09	0.19	-0.03	0.15
3.30	0.19	0.12	0.21	0.03	0.12
3.72	0.19	0.14	0.22	0.18	0.17
4.13	0.33	0.17	0.24	0.02	0.07

Table VIII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density	Cross-Talk (Cyan)	Cross-Talk (Yellow)
0.00	0.05	0.07	0.16		
0.67	0.05	0.07	0.16		
1.34	0.05	0.07	0.17		
2.01	0.05	0.07	0.18		
2.68	0.06	0.07	0.18		
3.36	0.06	0.08	0.18		
4.03	0.08	0.12	0.19		
4.70	0.08	0.24	0.22	0.16	0.17
5.37	0.10	0.38	0.25	0.14	0.11
6.04	0.16	0.63	0.33	0.18	0.12
6.71	0.20	0.91	0.42	0.16	0.13

Table IX

Voltage applied (V)	Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
7.5	639	0.06	0.26	0.73
7	557	0.06	0.23	0.70

5 This example shows that all three colors may be printed independently using a thermal head addressing the same side of an imaging member constructed as shown in Fig. 9.

EXAMPLE V

10 This example illustrates a three color imaging member such as illustrated in Fig. 10. The top image-forming layer produces a yellow color, using a unimolecular thermal reaction mechanism as described in U. S. Patent No. 5,350,870. The middle image-forming layer produces a magenta color, using an acid developer, an acid co-developer, and a magenta leuco dye. The bottom image-forming layer produces a cyan color, using an acid developer, and a cyan leuco dye. In between the magenta
15 and cyan layer, a thick clear poly(ethylene terephthalate) film base of approximately

102 micron thickness (Cronar 412) was used. Below the bottom cyan image-forming layer, a thick, opaque, white layer was used as a masking layer. The imaging member was addressed from the top (yellow and magenta) and the bottom (cyan). Because of the presence of the opaque layer, however, all three colors were visible only from the top. In this manner, a full-color image could be obtained.

A. The magenta image-forming layer was prepared as follows:

Dispersions of Leuco Dye I and Acid Developer I were prepared as described in Example I, part A. above.

A dispersion of Acid Developer III was prepared as described in Example II, part A. above.

The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated on a clear poly(ethylene terephthalate) film base of approximately 102 microns' thickness (Cronar 412) onto the gelatine-subcoated side, using a Meyer rod, and dried. The intended coating thickness was 3 microns.

Ingredient	% solids in dried film
Leuco Dye I	24.18%
Acid Developer I	47.49%
Acid Developer III	11.63%
Jonyl 138	16.16%
Zonyl FSN	0.54%

B. A thermally insulating interlayer was deposited onto the magenta imaging layer as described in Example II, part B. above.

C. A yellow image-forming layer was deposited on the thermally insulating layer as follows:

A dispersion of Leuco Dye III was prepared as described in Example II, part C. above. This dispersion was used to make the yellow coating fluid in proportions stated below. The yellow coating composition thus prepared was coated on the thermally insulating interlayer using a Meyer rod for an intended thickness of 3 microns, and was dried in air.

Ingredient	% solids in dried film
Leuco Dye III	70%
Genflo 3056	22.95%
Airvol 205	7%
Zonyl FSN	0.05%

D. A protective overcoat was deposited on the yellow image-forming layers as follows:

A slip overcoat was coated on the yellow dye layer. The overcoat was prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the yellow dye layer using a Meyer rod for an intended thickness of 1.0 microns, and was dried in air.

Ingredient	% solids in dried film
Glyoxal	8.39%
Hymicron ZK-349	31.77%
Klebosol 30V-25	23.77%
Zonyl FSA	0.92%
Zonyl FSN	3.22%
Airvol 540	31.93%

E. The cyan image-forming layer was prepared as follows:

Leuco Dye II was dispersed in an aqueous mixture comprising Airvol 205 (2.7% of total solids), Airvol 350 (6.3% of total solids), Triton X-100 (0.18% of total solids) and Aerosol-OT (0.9% of total solids) in deionized water, using an attriter equipped with glass beads and stirred for 18 hours at room temperature. The total solid content of the dispersion was 20%.

A dispersion of Acid Developer I was prepared as described in Example I, part A. above.

The above dispersions were used to make the cyan coating fluid in proportions stated below. The coating composition thus prepared was coated onto the opposite side of the clear poly(ethylene terephthalate) film base as coatings A-D, using a Meyer rod, and dried in air. The intended coating thickness was 2 microns.

Ingredient	% solids in dried film
Leuco Dye II	28.38%
Acid Developer I	41.62%
GenFlo 3056	22.90%
Airvol 205	7 %
Zonyl FSN	0.1%

F. The masking, opaque layer.

Titanium dioxide was dispersed in an aqueous mixture comprising Tamol 731 (3.86% of total solids), Ludox HS40 (3.85% of total solids) and a trace amount (750ppm) of Nipa Proxel in deionized water, using an attriter equipped with glass
5 beads and stirred for 18 hours at room temperature. The total solid content of the dispersion was 50.2%.

The above dispersion was used to make a coating fluid in proportions stated below. The coating composition thus prepared was coated on the cyan image-forming layer using a Meyer rod for an intended thickness of 15 micron, and was
10 dried in air.

Ingredient	% solids in dried film
Titanium dioxide	81.37%
Joncryl 138	18.08%
Zonyl FSN	0.54%

G. A protective overcoat was deposited on the opaque layer as described in part D. above.

The resulting imaging member was printed using a laboratory test-bed printer equipped with a thermal head, model KST-87-12MPC8 (Kyocera
15 Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan). The following printing parameters were used:

Printhead width: 3.41 inch
Pixels per inch: 300
Resistor size: 69.7 x 80 microns
20 Resistance: 3536 Ohm
Line Speed: 8 milliseconds per line
Print speed: 0.42 inches per second

Pressure: 1.5 - 2 lb/linear inch

Dot pattern: Rectangular grid.

The yellow layer was printed from the front side with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased
5 from zero to a maximum of 1.65 milliseconds (about 20.6% of the total line time) in twenty-one equal steps, while the voltage supplied to the print head was maintained at 29.0V.

A lower power/longer time condition was used to print the magenta layer, which was also addressed from the front side. The pulse width was increased from
10 zero to the 99.5% of 8 millisecond line time in twenty-one equal steps, while the voltage supplied to the print head was maintained at 16V.

The cyan layer was printed with a high power/short time condition from the backside (the side of the film base bearing the opaque layer). In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.65
15 milliseconds (about 20.6% of the total line time) in twenty-one equal steps, while the voltage supplied to the print head was maintained at 29.0V.

Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. The results are shown in Tables X, XI and XII. Table X shows the printing of the yellow layer as a function
20 of energy supplied by the thermal head. The magenta and cyan densities obtained are shown as well. Also included in Table X are the ratio between the yellow and the magenta density (Y/M) and the cross-talk. Similarly, Table XI shows the printing of the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the yellow densities is shown (M/Y) as
25 well as the cross-talk. In Table XII, printing of cyan layer as a function of the energy supplied by the thermal head is also listed. The ratio between the cyan and magenta densities is shown (C/M).

Table X

Energy Supplied (J/cm ²)	Yellow printed density	Magenta printed density	Cyan printed density	Y/M	Cross-Talk (Magenta)
0.00	0.11	0.11	0.08	1.00	
0.26	0.11	0.11	0.08	1.00	
0.52	0.11	0.11	0.08	1.00	
0.78	0.12	0.11	0.08	1.09	
1.04	0.11	0.11	0.08	1.00	
1.30	0.11	0.11	0.08	1.00	
1.56	0.12	0.11	0.08	1.09	
1.82	0.12	0.11	0.08	1.09	
2.08	0.13	0.11	0.08	1.18	
2.34	0.15	0.11	0.08	1.36	
2.60	0.21	0.12	0.08	1.75	-0.01
2.86	0.28	0.12	0.08	2.33	-0.05
3.12	0.36	0.13	0.08	2.77	-0.03
3.38	0.46	0.15	0.08	3.07	0.01
3.64	0.63	0.17	0.08	3.71	0.01
3.90	0.79	0.20	0.08	3.95	0.03
4.16	0.98	0.24	0.08	4.08	0.05
4.41	1.12	0.27	0.08	4.15	0.06
4.67	1.24	0.30	0.09	4.13	0.06
4.93	1.36	0.33	0.09	4.12	0.07
5.19	1.44	0.36	0.09	4.00	0.08

Table XI

Energy Supplied (J/cm ²)	Magenta printed density	Yellow printed density	Cyan printed density	M/Y	Cross-Talk (Yellow)
0.00	0.11	0.11	0.07	1.00	
0.38	0.11	0.11	0.08	1.00	
0.76	0.11	0.11	0.07	1.00	
1.15	0.11	0.11	0.08	1.00	
1.53	0.11	0.11	0.08	1.00	
1.91	0.11	0.11	0.08	1.00	
2.29	0.11	0.11	0.08	1.00	
2.67	0.11	0.11	0.07	1.00	
3.05	0.11	0.11	0.07	1.00	
3.44	0.11	0.12	0.07	0.92	
3.82	0.11	0.12	0.07	0.92	
4.20	0.12	0.13	0.07	0.92	
4.58	0.13	0.14	0.07	0.93	
4.96	0.17	0.16	0.07	1.06	
5.35	0.24	0.19	0.08	1.26	0.47
5.73	0.39	0.25	0.09	1.56	0.34
6.11	0.60	0.34	0.10	1.76	0.31
6.49	0.86	0.44	0.12	1.95	0.28
6.87	1.16	0.55	0.13	2.11	0.25
7.26	1.50	0.71	0.15	2.11	0.27
7.64	1.54	0.81	0.16	1.90	0.33

Table XII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density	C/M
0.00	0.07	0.11	0.11	0.64
0.26	0.07	0.11	0.11	0.64
0.52	0.07	0.11	0.11	0.64
0.78	0.07	0.11	0.11	0.64
1.04	0.07	0.11	0.11	0.64
1.30	0.07	0.11	0.11	0.64
1.56	0.07	0.11	0.11	0.64
1.82	0.07	0.11	0.11	0.64
2.08	0.07	0.11	0.11	0.64
2.34	0.07	0.11	0.11	0.64
2.60	0.08	0.11	0.11	0.73
2.86	0.10	0.11	0.11	0.91
3.12	0.16	0.13	0.12	1.23
3.38	0.24	0.15	0.13	1.60
3.64	0.33	0.17	0.14	1.94
3.90	0.43	0.21	0.15	2.05
4.16	0.57	0.26	0.18	2.19
4.41	0.90	0.42	0.27	2.14
4.67	1.09	0.53	0.33	2.06
4.93	1.06	0.52	0.33	2.04
5.19	1.03	0.51	0.32	2.02

EXAMPLE VI

This example illustrates a three color imaging member such as illustrated in Fig. 10. The top image-forming layer produces a cyan color, the middle image-forming layer produces a magenta color, and the bottom image-forming layer produces a yellow color. All three layers use an acid developer or developers, and a leuco dye. In between the magenta and yellow layers, a thick clear poly(ethylene terephthalate) film base of approximately 102 micron thickness (Cronar 412) was used. Below the bottom yellow image-forming layer, a thick, opaque, white layer was used as a masking layer. The imaging member was addressed from the top (cyan and magenta) and the bottom (yellow). Because of the presence of the opaque layer, however, all three colors were visible only from the top. In this manner, a full-color image could be obtained.

A. The magenta color-forming layer was prepared as follows:

Dispersions of Leuco Dye I and Acid Developer I were prepared as described in Example IV, part C above. A dispersion of Acid Developer II was prepared as described in Example II, part A above.

5 The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated onto Cronar 412, and dried. The intended coating coverage was 2.0 g/m².

Ingredient	% solids in dried film
Leuco Dye I	24.18%
Acid Developer I	47.50%
Joncryl 138	16.16%
Acid Developer II	11.63%
Zonyl FSN	0.54%

B. A thermally insulating interlayer was deposited onto the magenta imaging layer as follows:

10 A coating fluid for the interlayer was prepared in proportions stated below. The image interlayer coating composition thus prepared was coated on the magenta imaging layer in three passes, for an intended coverage of 13.4 g/m².

Ingredient	% solids in dried film
Glascol C44	99.50%
Zonyl FSA	0.50%

C. Cyan image-forming layers C1 – C3 were deposited on the thermally insulating layer as follows:

15 C1 Cyan developer layer.

A dispersion of Acid Developer III was prepared as described in Example IV, part E1 above.

20 The above dispersion was used to make the cyan developer coating fluid in proportions stated below. The cyan developer coating composition thus prepared was coated on top of the thermally-insulating interlayer for an intended thickness of 2.1 g/m², and was dried.

Ingredient	% solids in dried film
Joncryn 138	10.0%
Acid Developer III	89.5%
Zonyl FSN	0.50%

C2 Cyan interlayer.

A cyan interlayer coating fluid was prepared in proportions stated below. The cyan interlayer coating composition thus prepared was coated on top of the cyan developer layer for an intended thickness of 1.0 g/m².

Ingredient	% solids in dried film
Airvol 205	99.00%
Zonyl FSN	1.00%

5 C3 Cyan dye layer.

Leuco dye II was dispersed as described in Example IV, part E3 above.

The above dispersion was used to make the cyan coating fluid in proportions stated below. The cyan coating composition thus prepared was coated on the cyan interlayer for an intended coverage of 0.65 g/m².

Ingredient	% solids in dried film
Leuco Dye II	59.30%
Joncryn 138	39.37%
Zonyl FSN	1.33%

10 D. A protective overcoat was deposited on the cyan image-forming layers as follows:

A slip overcoat was coated on the cyan dye layer. The overcoat was prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the cyan dye layer for an intended coverage of 1.1 g/m².

Ingredient	% solids in dried film
Hymicron ZK-349	31.77%
Klebosol 30V-25	23.77%
Airvol 540	31.93%
Glyoxal	8.39%
Zonyl FSA	0.92%
Zonyl FSN	3.22%

E. A yellow image-forming layer was deposited onto the reverse of the clear substrate using the procedure described in Example IV, part A above, except that the dried coverage was 1.94 g/m².

F. A white, opaque layer was deposited onto the yellow color-forming layer as follows:

A dispersion of titanium dioxide was prepared as described in Example V, part F. above.

A coating fluid was prepared from the dispersion so formed in proportions stated below. The coating composition thus prepared was coated on top of the yellow color-forming layer for an intended coverage of 10.76 g/m².

Ingredient	% solids in dried film
Titanium dioxide	89.70%
Joncryl 138	9.97%
Zonyl FSN	0.33%

G. A protective overcoat was deposited on the opaque layer as described in part D. above.

The resulting imaging member was printed using a laboratory test-bed printer equipped with a thermal head, model KST-87-12MPC8 (Kyocera Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan). The following printing parameters were used:

Printhead width: 3.41 inch
Pixels per inch: 300
Resistor size: 69.7 x 80 microns
Resistance: 3536 Ohm
Line Speed: 8 milliseconds per line
Print speed: 0.42 inches per second
Pressure: 1.5 - 2 lb/linear inch
Dot pattern: Rectangular grid.

The cyan layer was printed from the front side with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.25 milliseconds (about 16.4% of the total line time) in

twenty-one equal steps, while the voltage supplied to the print head was maintained at 29.0V.

5 A lower power/longer time condition was used to print the magenta layer, which was also addressed from the front side. The pulse width was increased from zero to the 99.5% of 8 millisecond line time in twenty-one equal steps, while the voltage supplied to the print head was maintained at 14.5V.

10 The yellow layer was printed with a lower power/longer time condition from the backside (the side of the film base bearing the opaque layer). The pulse width was increased from zero to the 99.5% of 8 millisecond line time in twenty-one equal steps, while the voltage supplied to the print head was maintained at 14.5V.

15 Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. The results are shown in Tables XIII, XIV and XV. Table XIII shows the printing of the cyan layer as a function of energy supplied by the thermal head. The magenta and yellow densities obtained are shown as well. Also included in Table XIII are the ratio between the cyan and the magenta density (C/M) and the cross-talk. Similarly, Table XIV shows the printing of the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the cyan densities is shown (M/C) as well as the cross-talk. In Table XV, printing of yellow layer as a function of the energy supplied by the thermal head is also listed. The ratio between the yellow and magenta densities is shown (Y/M).

20

Table XIII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density	C/M	Cross-Talk (Magenta)
1.57	0.07	0.10	0.23	0.70	
1.83	0.08	0.10	0.23	0.80	
2.09	0.08	0.11	0.25	0.73	
2.34	0.08	0.10	0.23	0.80	
2.60	0.11	0.11	0.23	1.00	
2.85	0.12	0.12	0.23	1.00	
3.11	0.16	0.13	0.24	1.23	-0.01
3.36	0.20	0.14	0.25	1.43	-0.04
3.62	0.26	0.16	0.26	1.63	-0.03
3.87	0.28	0.17	0.27	1.65	-0.01
4.13	0.36	0.20	0.28	1.80	0.00

Table XIV

Energy Supplied (J/cm ²)	Magenta printed density	Cyan printed density	Yellow printed density	M/C	Cross-Talk (Cyan)
3.14	0.10	0.07	0.20	1.43	
3.45	0.11	0.09	0.22	1.22	
3.76	0.11	0.09	0.22	1.22	
4.08	0.12	0.10	0.22	1.20	
4.39	0.13	0.10	0.21	1.30	
4.70	0.16	0.11	0.23	1.45	
5.02	0.21	0.11	0.24	1.91	0.39
5.33	0.30	0.14	0.24	2.14	0.36
5.65	0.43	0.16	0.26	2.69	0.27
5.96	0.57	0.17	0.29	3.35	0.20
6.27	0.60	0.18	0.29	3.33	0.20

Table XV

Energy Supplied (J/cm ²)	Yellow printed density	Magenta printed density	Cyan printed density	Y/M
0.00	0.23	0.10	0.07	2.30
0.63	0.23	0.10	0.07	2.30
1.25	0.24	0.10	0.08	2.40
1.88	0.22	0.10	0.08	2.20
2.51	0.22	0.10	0.07	2.20
3.14	0.23	0.10	0.08	2.30
3.76	0.32	0.10	0.07	3.20
4.39	0.57	0.12	0.07	4.75
5.02	0.85	0.18	0.07	4.72
5.65	0.95	0.25	0.07	3.80
6.27	0.98	0.33	0.08	2.97

EXAMPLE VII

This example illustrates the preparation of the zinc salt of 3-methyl-5-n-octylsalicylic acid.

5 Preparation of methyl 3-methyl-5-n-octanoyl salicylate:

Aluminum chloride (98 g) was suspended in methylene chloride (150 mL) in a 1L flask and the mixture was cooled to 5° C. in an ice bath. To the stirred mixture was added methyl 3-methylsalicylate (50 g) and octanoyl chloride (98 g) in 150 mL of methylene chloride over a 1hr period. The reaction was stirred for an additional
 10 30 min. at 5° C and then at 3 hrs at room temperature. The reaction was poured into 500g of ice containing 50mL of concentrated hydrochloric acid. The organic layer was separated and the aqueous layer extracted twice with 50mL of methylene chloride. The methylene chloride was washed with a saturated aqueous solution of sodium bicarbonate, dried with magnesium sulfate, filtered, and evaporated to an oil
 15 which solidified to 90g of tan crystals. ¹H and ¹³C NMR spectra were consistent with expected product.

Preparation of 3-methyl-5-n-octanoyl salicylic acid:

Methyl 3-methyl-5-n-octanoyl salicylate (prepared as described above, 90 g) was dissolved in 200mL of ethanol and 350mL of water. To this solution was added
 20 100g of a 50% aqueous solution of sodium hydroxide and the solution was than stirred at 85° C for 6hrs. The reaction was cooled in an ice bath and a 50% aqueous

solution of hydrochloric acid was slowly added until a pH of 1 was attained. The precipitate was filtered, washed with water (5x50mL) and dried under reduced pressure at 45° C for 6hrs. to give 80g of pale tan product. ¹H and ¹³C NMR spectra were consistent with expected product.

5 Preparation of 3-methyl-5-n-octyl salicylic acid:

16g of mercury(II) chloride was dissolved in 8mL of concentrated hydrochloric acid and 200 mL of water in a 1L flask. 165g Mossy zinc was shaken with this solution. The water was decanted off and to the zinc was added 240mL of concentrated hydrochloric acid, 100mL of water and 3-methyl-5-n-octanoyl salicylic acid (prepared as described above, 80 g). The mixture was refluxed with stirring for 10 24 hrs. with an additional 50mL of concentrated hydrochloric acid being added every 6hrs (3 times). The reaction was decanted hot from the zinc and cooled to solidify the product. The product was collected by filtration, washed with (2x 100mL water) and dissolved in 300mL hot ethanol. 50mL of water was added and 15 the solution was refrigerated to give white crystals. The solid was filtered, washed (3x 100mL water) and dried under reduced pressure at 45° C for 8hrs to give 65g of product. ¹H and ¹³C NMR spectra were consistent with expected product.

Preparation of 3-methyl-5-n-octyl salicylic acid zinc salt:

3-Methyl-5-n-octyl salicylic acid (prepared as described above, 48 g) was 20 added with stirring to a solution of 14.5g of a 50% aqueous solution of sodium hydroxide and 200mL water in a 4L beaker. To this was added 1L of water and the solution was heated to 65° C. To the hot solution was then added with stirring 24.5g of zinc chloride in 40ml of water. A gummy solid precipitated. The solution decanted and the remaining solid was dissolved in 300mL hot 95% ethanol. The hot 25 solution was diluted with 500ml of water and refrigerated. The product was filtered and washed (3x 500mL water) to give 53g of off-white solid.

EXAMPLE VIII

30 This example illustrates a three color imaging member with an overcoat layer deposited on each side, and a method for writing multiple colors on this member in a

single pass using two thermal print heads. The top color-forming layer produces a yellow color, using a unimolecular thermal reaction mechanism as described in U. S. Patent No. 5,350,870. The middle color-forming layer produces a magenta color, using an acid developer, an acid co-developer, and a magenta leuco dye. The bottom color-forming layer produces a cyan color, using an acid developer, and a cyan leuco dye. In between the magenta and cyan layer, a thick clear poly(ethylene terephthalate) film base of approximately 102 micron thickness (Cronar 412) was used. Below the bottom cyan image-forming layer, a thick, opaque, white layer was used as a masking layer. The imaging member was addressed from the top (yellow and magenta) and the bottom (cyan). Because of the presence of the opaque layer, however, all three colors were visible only from the top. In this manner, a full-color image could be obtained.

A. The magenta image-forming layer was prepared as follows:
Dispersions of Leuco Dye I and Acid Developer I were prepared as described in Example I, part A. above.

A dispersion of Acid Developer III was prepared as described in Example II, part A. above.

The above dispersions were used to make the magenta coating fluid in proportions stated below. The coating composition thus prepared was coated on a clear poly(ethylene terephthalate) film base of approximately 102 microns' thickness (Cronar 412) onto the gelatin-subcoated side, using a Meyer rod, and dried. The intended coating thickness was 3.06 microns.

Ingredient	% solids in dried film
Leuco Dye I	12.08%
Acid Developer I	28.70%
Acid Developer II	15.14%
Genflo 3056	37.38%
Airvol 205	6.38%
Zonyl FSN	0.32%

B. A thermally insulating interlayer was deposited onto the magenta imaging layer as follows:

B1. A coating fluid for the interlayer was prepared in the proportions stated below. The image interlayer coating composition thus prepared was coated on the imaging layer using a Meyer rod for an intended thickness of 6.85 microns, and was dried in air.

Ingredient	% solids in dried film
Glascol C44	99.78%
Zonyl FSN	0.22%

B2. A second insulating interlayer of the same description was then coated on the first interlayer and dried.

B3. Finally, a third insulating interlayer of the same description was coated on the second interlayer and dried. The combination of the three insulating interlayers comprised an insulating layer with an intended total thickness of 20.55 microns.

C. A yellow image-forming layer was deposited on the third thermally insulating layer as follows:

A dispersion of Leuco Dye III was prepared as described in Example II, part C. above. This dispersion was used to make the yellow coating fluid in proportions stated below. The yellow coating composition thus prepared was coated on the thermally insulating interlayer using a Meyer rod for an intended thickness of 3.21 microns, and was dried in air.

Ingredient	% solids in dried film
Leuco Dye III	49.42%
Airvol 205	11.68%
Genflo 3056	38.00%
Zonyl FSN	0.90%

D. A protective overcoat was deposited on the yellow image-forming layers as follows:

A slip overcoat was coated on the yellow dye layer. The overcoat was prepared in proportions stated below. The overcoat coating composition thus prepared was coated on the yellow dye layer using a Meyer rod for an intended thickness of 1.46 microns, and was dried in air.

Ingredient	% solids in dried film
Glyoxal	8.54%
Hymicron ZK-349	31.95%
Klebosol 30V-25	23.89%
Zonyl FSA	0.98%
Zonyl FSN	2.44%
Airvol 540	32.20%

E. The cyan image-forming layer was prepared as follows:

Leuco Dye II was dispersed in an aqueous mixture comprising Airvol 205 (2.7% of total solids), Airvol 350 (6.3% of total solids), Triton X-100 (0.18% of total solids) and Aerosol-OT (0.9% of total solids) in deionized water, using an attriter equipped with glass beads and stirred for 18 hours at room temperature. The total solid content of the dispersion was 20%.

A dispersion of Acid Developer I was prepared as described in Example I, part A. above.

The above dispersions were used to make the cyan coating fluid in proportions stated below. The coating composition thus prepared was coated onto the opposite side of the clear poly(ethylene terephthalate) film base as coatings A-D, using a Meyer rod, and dried in air. The intended coating thickness was 3.01 microns.

Ingredient	% solids in dried film
Leuco Dye II	18.94%
Acid Developer I	51.08%
GenFlo 3056	22.86%
Airvol 205	7.01 %
Zonyl FSN	0.10%

F. The masking, opaque layer.

Titanium dioxide was dispersed in an aqueous mixture comprising Tamol 731 (3.86% of total solids), Ludox HS40 (3.85% of total solids) and a trace amount (750ppm) of Nipa Proxel in deionized water, using an attriter equipped with glass beads and stirred for 18 hours at room temperature. The total solid content of the dispersion was 50.2%.

The above dispersion was used to make a coating fluid in proportions stated below. The coating composition thus prepared was coated on the cyan image-forming layer using a Meyer rod for an intended thickness of 15 micron, and was dried in air.

Ingredient	% solids in dried film
Titanium dioxide	88.61%
Airvol 205	11.08%
Zonyl FSN	0.32%

- 5 G. A protective overcoat was deposited on the opaque layer as described in part D. above.

The resulting imaging member was printed using a laboratory test-bed printer equipped with two thermal heads, model KYT-106-12PAN13 (Kyocera Corporation, 6 Takedatobadono-cho, Fushimi-ku, Kyoto, Japan). The following printing parameters were used:

Printhead width: 4.16 inch
Pixels per inch: 300
Resistor size: 70 x 80 microns
15 Resistance: 3900 Ohm
Line Speed: 10.7 milliseconds per line
Print speed: 0.31 inches per second
Pressure: 1.5 - 2 lb/linear inch
Dot pattern: Rectangular grid.

20 The yellow layer was printed from the front side with a high power/short time condition. In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 1.99 milliseconds (about 18.2% of the total line time) in ten equal steps, while the voltage supplied to the print head was maintained at 26.5V. Within this pulse width there were 120 subintervals, and each had a duty
25 cycle of 95%.

A lower power/longer time condition was used to print the magenta layer, which was also addressed from the front side. The pulse width was increased from

zero to a maximum of 8.5 milliseconds (about 79% of the total line time) in 10 equal steps, while the voltage supplied to the print head was maintained at 26.5V. Within this pulse width, there were 525 subintervals, and each had a duty cycle of 30%.

Unlike previous examples, the yellow pulses and magenta pulses were
5 interleaved, and were supplied by a single print head in a single pass, so that a single printhead was printing two colors synchronously. The selection of high power or low power was made by alternating between the 95% duty cycle used for printing yellow and the 30% duty cycle used for printing magenta. The print head voltage was constant at 26.5V.

10 The cyan layer was printed with a low-power, long-time condition from the backside (the side of the film base bearing the opaque TiO_2 layer). In order to obtain gradations of color, the pulse width was increased from zero to a maximum of 10.5 milliseconds (about 98% of the total line time) in 10 equal steps, while the voltage supplied to the print head was maintained at 21.0V.

15 In addition to printing gradations of color for each of the three dye layers, gradations of combined pairs of the colors, and of the combination of all three colors, were printed.

Following printing, the reflection density in each of the printed areas was measured using a Gretag Macbeth spectrophotometer. Results for writing on the
20 yellow, magenta and cyan layers are shown in Tables XVI, XVII and XVIII.

Table XVI shows the printing of the cyan layer as a function of energy supplied by the thermal head. The magenta and yellow densities obtained are shown as well. Similarly, Table XVII shows the printing of the magenta layer as a function of the energy supplied by the thermal head. The ratio between the magenta and the
25 yellow densities is also shown (M/Y) as well as the cross-talk. In Table XVIII, printing of yellow layer as a function of the energy supplied by the thermal head is also listed. The ratio between the yellow and magenta densities is shown (Y/M) as well as the cross-talk.

Table XVI

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
1.79	0.10	0.12	0.20
2.07	0.11	0.12	0.20
2.35	0.11	0.12	0.19
2.63	0.12	0.13	0.19
2.92	0.17	0.13	0.20
3.20	0.25	0.15	0.20
3.48	0.34	0.18	0.22
3.76	0.56	0.25	0.25
4.05	0.82	0.35	0.29
4.33	1.07	0.43	0.33
4.61	1.17	0.45	0.34

Table XVII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density	M/Y	Cross-Talk Yellow
3.07	0.11	0.13	0.20	0.65	
3.40	0.10	0.13	0.20	0.65	
3.74	0.10	0.13	0.20	0.65	
4.08	0.10	0.14	0.22	0.64	
4.42	0.10	0.16	0.22	0.73	
4.75	0.10	0.21	0.24	0.88	
5.09	0.11	0.33	0.27	1.22	0.18
5.43	0.11	0.53	0.31	1.71	0.11
5.77	0.13	0.80	0.38	2.10	0.10
6.10	0.14	0.97	0.43	2.25	0.10
6.45	0.14	1.02	0.45	2.27	0.11

Table XVIII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density	Y/M	Cross-Talk Magenta
1.82	0.11	0.13	0.20	1.53	
2.07	0.11	0.13	0.22	1.69	
2.33	0.11	0.13	0.27	2.08	
2.58	0.10	0.13	0.31	2.38	
2.84	0.11	0.14	0.36	2.57	
3.09	0.10	0.15	0.48	3.20	
3.35	0.11	0.17	0.59	3.47	0.00
3.60	0.11	0.19	0.71	3.74	0.01
3.86	0.11	0.20	0.76	3.80	0.02
4.11	0.11	0.21	0.88	4.19	0.01
4.37	0.11	0.21	0.84	4.00	0.02

The results obtained by writing on combinations of two color layers are shown in Tables XIX, XX and XXI. Table XIX illustrates the result of printing simultaneously on the yellow and magenta layers with a single thermal print head. The resulting print is red in color. Table XX shows the result of printing simultaneously on the cyan and yellow layers, giving a green print, and Table XXI shows the result of printing on the cyan and magenta layers to give a blue print.

Table XIX

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
4.89	0.10	0.12	0.20
5.47	0.11	0.14	0.23
6.08	0.11	0.17	0.28
6.66	0.11	0.27	0.38
7.26	0.12	0.40	0.50
7.84	0.13	0.80	0.65
8.45	0.15	1.20	0.84
9.03	0.18	1.60	1.11
9.63	0.19	1.71	1.26
10.21	0.19	1.69	1.39
10.82	0.20	1.62	1.42

Table XX

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
3.61	0.11	0.13	0.20
4.14	0.11	0.13	0.20
4.69	0.12	0.13	0.22
5.21	0.13	0.14	0.27
5.76	0.17	0.15	0.32
6.29	0.31	0.19	0.43
6.84	0.46	0.26	0.55
7.36	0.67	0.33	0.57
7.91	0.92	0.43	0.67
8.44	1.23	0.54	0.84
8.99	1.36	0.58	0.93

Table XXI

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
4.86	0.11	0.12	0.19
5.47	0.11	0.13	0.24
6.10	0.12	0.13	0.20
6.71	0.13	0.15	0.21
7.34	0.15	0.17	0.22
7.95	0.32	0.26	0.25
8.58	0.51	0.42	0.31
9.19	0.69	0.76	0.39
9.82	0.88	1.01	0.47
10.43	1.40	1.27	0.59
11.06	1.49	1.31	0.61

Table XXII presents the color densities resulting from printing on all three color layers in a single pass. The resulting print is black.

Table XXII

Energy Supplied (J/cm ²)	Cyan printed density	Magenta printed density	Yellow printed density
6.68	0.11	0.13	0.20
7.54	0.11	0.14	0.24
8.43	0.11	0.17	0.29
9.29	0.11	0.23	0.37
10.18	0.18	0.43	0.43
11.04	0.29	0.81	0.71
11.93	0.41	1.21	0.94
12.79	0.64	1.59	1.12
13.68	0.89	1.81	1.38
14.54	1.17	1.79	1.46
15.43	1.29	1.71	1.55

Although the invention has been described in detail with respect to various preferred embodiments, it is not intended to be limited thereto, but rather those skilled in the art will recognize that variations and modifications are possible which are within the spirit of the invention and the scope of the appended claims.